

(FILE 'HOME' ENTERED AT 11:18:52 ON 27 JUN 2005)

FILE 'CAPLUS' ENTERED AT 11:19:07 ON 27 JUN 2005

L1                   STRUCTURE UPLOADED  
                  S L1

FILE 'REGISTRY' ENTERED AT 11:19:31 ON 27 JUN 2005

L2                   50 S L1

FILE 'CAPLUS' ENTERED AT 11:19:32 ON 27 JUN 2005

L3                   3 S L2  
                  S L1

FILE 'REGISTRY' ENTERED AT 11:20:12 ON 27 JUN 2005

L4                   21641 S L1 FULL

FILE 'CAPLUS' ENTERED AT 11:20:13 ON 27 JUN 2005

L5                   13121 S L4 FULL  
L6                   10617 S L5 AND PY<1999  
L7                   927 S L6 AND (ESTER OR AMIDE)  
L8                   316 S L7 AND QUATERN?  
L9                   17 S L8 AND DICARBOXYLIC ACID

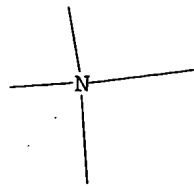
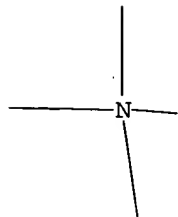
=>

L1 STRUCTURE UPLOADED

=> d

L1 HAS NO ANSWERS

L1 STR



Structure attributes must be viewed using STN Express query preparation.

L9 ANSWER 1 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1989:205078 CAPLUS  
 DOCUMENT NUMBER: 110:205078  
 TITLE: Relations between structure, hydrolysis rate and activity of **dicarboxylic acid** esters  
 AUTHOR(S): Kharkevich, D. A.; Skoldinov, A. P.; Lemina, E. Yu.; Igumnova, N. D.  
 CORPORATE SOURCE: Dep. Pharmacol., First Med. Inst., Moscow, 119881, USSR  
 SOURCE: Farmakologiya i Toksikologiya (Moscow) (1989), 52(2), 34-7  
 CODEN: FATOAO; ISSN: 0014-8318  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Russian

AB The kinetics of enzymic cholinesterase hydrolysis of **dicarboxylic acid** esters [MeN(R)-(CH<sub>2</sub>)<sub>n</sub>-O<sub>2</sub>C-(CH<sub>2</sub>)<sub>m</sub>-CO<sub>2</sub>-(CH<sub>2</sub>)<sub>n</sub>-(R)NMe + 2 MeI or 2 HCl, R = Me, 1-adamantyl; m = 1,2,4,6,8; n = 2,4] with neuromuscular-blocking activity was studied in vitro. The maximum hydrolysis rate was shown to increase on elongation of the distance between **ester** groups, both in the compds. containing a hydrophobic adamantyl radical attached to **quaternary** nitrogen, and in bis esters not containing adamantyl radicals. The comparison of neuromuscular-blocking activity in vivo, enzymic hydrolysis rates, and activity on isolated skeletal muscle demonstrated that in vivo activity is more strongly correlated with the maximum hydrolysis rate of the compds. than with activity in isolated skeletal muscle.

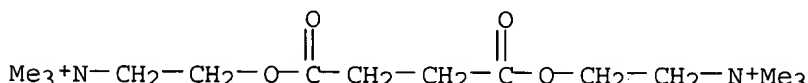
IT 541-19-5 1807-06-3 71677-28-6

71677-29-7 71677-30-0

RL: RCT (Reactant); RACT (Reactant or reagent)  
 (hydrolysis of, rate of, neuromuscular-blocking activity and structure in relation to)

RN 541-19-5 CAPLUS

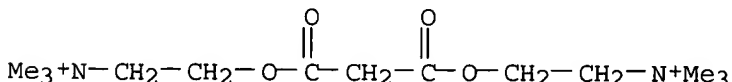
CN Ethanaminium, 2,2'-(1,4-dioxo-1,4-butanediyl)bis(oxy)bis[N,N,N-trimethyl-, diiodide (9CI) (CA INDEX NAME)



● 2 I-

RN 1807-06-3 CAPLUS

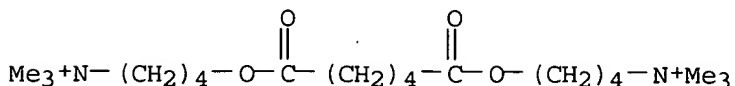
CN Ethanaminium, 2,2'-(1,3-dioxo-1,3-propanediyl)bis(oxy)bis[N,N,N-trimethyl-, diiodide (9CI) (CA INDEX NAME)



● 2 I-

RN 71677-28-6 CAPLUS

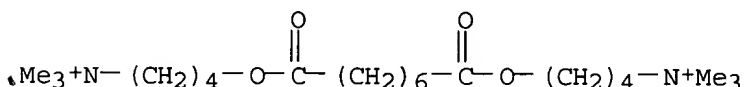
CN 1-Butanaminium, 4,4'-(1,6-dioxo-1,6-hexanediyi)bis(oxy)bis[N,N,N-trimethyl-, diiodide (9CI) (CA INDEX NAME)



●2 I-

RN 71677-29-7 CAPLUS

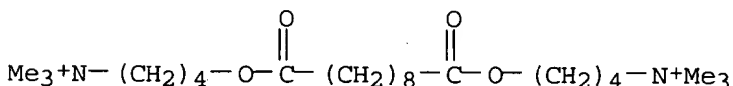
CN 1-Butanaminium, 4,4'-(1,8-dioxo-1,8-octanediyi)bis(oxy)bis[N,N,N-trimethyl-, diiodide (9CI) (CA INDEX NAME)



●2 I-

RN 71677-30-0 CAPLUS

CN 1-Butanaminium, 4,4'-(1,10-dioxo-1,10-decanediyl)bis(oxy)bis[N,N,N-trimethyl-, diiodide (9CI) (CA INDEX NAME)



●2 I-

L9 ANSWER 2 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1974:520077 CAPLUS

DOCUMENT NUMBER: 81:120077

TITLE: Bis-quaternary ammonium salts containing an adamantlyl radical

AUTHOR(S): Klimova, N. V.; Lavrova, L. N.; Skoldinov, A. P.; Kharkevich, D. A.; Shmar'yan, M. I.

CORPORATE SOURCE: Inst. Farmakol., Moscow, USSR

SOURCE: Khimiko-Farmatsevticheskii Zhurnal (1974), 8 (7), 3-5

DOCUMENT TYPE: CODEN: KHFZAN; ISSN: 0023-1134

LANGUAGE: Journal

RUSSIAN

GI For diagram(s), see printed CA Issue.

AB Reaction of  $\text{I}(\text{CH}_2)_n\text{I}$  ( $n = 5, 6, 7$ ) with  $\text{RNMe}_2$  ( $\text{R} = 1\text{-adamantyl}$ ) gave 50-62.5% the corresponding  $\text{Me}_2\text{N}^+\text{R}(\text{CH}_2)_n\text{N}^+-\text{Me}_2\text{R}\cdot 2\text{I}^-$  (I). I ( $n = 6$ ) was a ganglion-blocking agent at 0.12-0.2 mg/kg in cats (hexonium = 40 mg/kg). Transesterification of di-Me 1,3-adamantanedicarboxylate with  $\text{HO}(\text{CH}_2)_2\text{NMe}_2$ , followed by reaction of the resulting ester with  $\text{MeI}$  gave the bisammonium compound II. Reaction of 1,1'-diadamantyl-3,3'-dicarboxylic acid chloride with 1-methylpiperazine

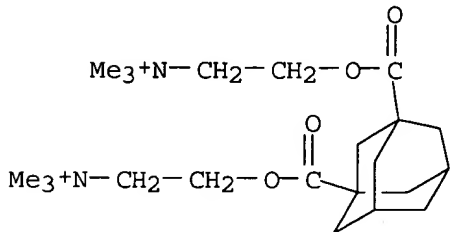
followed by MeI gave the bis(piperazinium salt III. III was an effective ganglion-blocking agent at 4-5 mg/kg in cats.

IT 51896-22-1P

RL: SPN (Synthetic preparation); PREP (Preparation)  
(preparation and ganglion blocking activity of)

RN 51896-22-1 CAPLUS

CN Ethanaminium, 2,2'-[tricyclo[3.3.1.13,7]decane-1,3-diy]bis[N,N,N-trimethyl-, diiodide (9CI) (CA INDEX NAME)



●2 I-

L9 ANSWER 3 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1973:92439 CAPLUS

DOCUMENT NUMBER: 78:92439

TITLE: Cyclobutane analogs of acetyl- $\gamma$ -homocholine

AUTHOR(S): Cannon, Joseph G.; Lin, Youlin; Long, John Paul

CORPORATE SOURCE: Coll. Pharm., Univ. Iowa, Iowa City, IA, USA

SOURCE: Journal of Medicinal Chemistry (1973),

16(1), 27-9

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE: Journal

LANGUAGE: English

AB Cis-(2-acetoxycyclobutylmethyl)trimethylammonium iodide (cis-I) [38868-89-2] and trans-(2-acetoxycyclobutylmethyl)trimethylammonium iodide (trans-I) [38868-90-5] had 40,000-fold and 5000-fold lower muscarinic activity, resp., than acetylcholine [51-84-3] in the superfused guinea pig ileum in vitro. The effects of both were blocked by atropine [51-55-8] but not by hexamethonium [60-26-4]. To synthesize cis-I, cis-cyclobutane-1,2-dicarboxylic acid mono-Me ester [31420-52-7] was converted with SOC12 to the mono-Me ester monoacyl chloride, with Me2CD to the mono-Me ester Me ketone, and with m-chloroperbenzoic acid to Me cis-2-acetoxycyclobutanecarboxylate [38868-92-7]. Aminolysis with NHMe2 gave a mixture of products which was reduced with LiAlH4 to cis-2-dimethylaminomethylcyclobutanol [38868-93-8]. This was quaternized with MeI and acetylated to yield cis-I. Trans-I was synthesized similarly.

IT 60-26-4

RL: BIOL (Biological study)  
(parasympathomimetic effects of acetyl homocholine derivs. in response to)

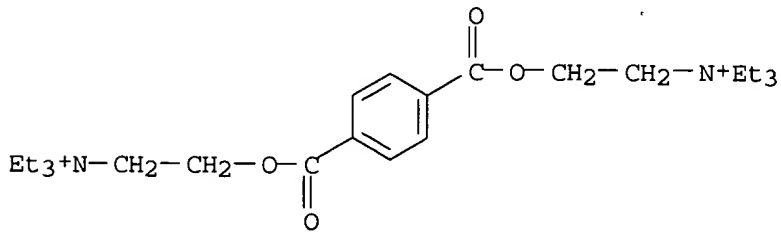
RN 60-26-4 CAPLUS

CN 1,6-Hexanediaminium, N,N,N,N',N',N'-hexamethyl- (9CI) (CA INDEX NAME)

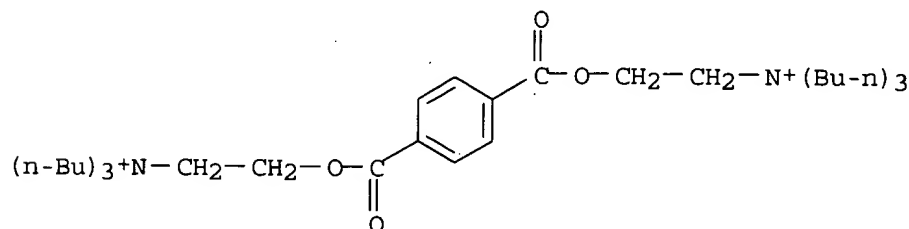
Me3+N-(CH2)6-N+Me3

L9 ANSWER 4 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1972:140248 CAPLUS  
 DOCUMENT NUMBER: 76:140248  
 TITLE: Glycol esters  
 INVENTOR(S): Kamatani, Hiroyoshi  
 PATENT ASSIGNEE(S): Toyo Spinning Co., Ltd.  
 SOURCE: Jpn. Tokkyo Koho  
 CODEN: JAXXAD  
 DOCUMENT TYPE: Patent  
 LANGUAGE: Japanese  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	JP 47002623	B4	19720125	JP	19680924 <--
AB	An aromatic dicarboxylic acid was made to react with an alkylene oxide, with use of a quaternary ammonium aromatic carboxylate having at least 1 N-(2-hydroxyalkyl) group as a catalyst. E.g., terephthalic acid was heated 100 min at 120° with ethylene oxide in xylene using 0.3 molar % bis(2-hydroxyethyltriethylammonium) terephthalate as a catalyst to give 75% ester. Examples of other catalysts used are bis(2-hydroxyethyltributylammonium) terephthalate and mono-(2-hydroxyethyldiethylcyclohexylammonium) terephthalate.				
IT	35719-59-6 35719-60-9 RL: CAT (Catalyst use); USES (Uses) (catalysts, for esterification of terephthalic acid by ethylene oxide)				
RN	35719-59-6 CAPLUS				
CN	Ethanaminium, 2,2'-[1,4-phenylenebis(carbonyloxy)]bis[N,N,N-triethyl- (9CI) (CA INDEX NAME)				



RN 35719-60-9 CAPLUS  
 CN 1-Butanaminium, N,N'-[1,4-phenylenebis(carbonyloxy-2,1-ethanediyl)]bis[N,N-dibutyl- (9CI) (CA INDEX NAME)]



L9 ANSWER 5 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1968:486735 CAPLUS  
 DOCUMENT NUMBER: 69:86735  
 TITLE: Acetylcholine. XII. 3,4-Diphenylthiophene-2.5-

**dicarboxylic acid bis  
[( $\beta$ -diethylamino)ethyl ester  
methiodide], a curarelike muscle-relaxant  
ester**

AUTHOR(S): Dann, O.; Bamberg, K. J.; Sucker, H.  
CORPORATE SOURCE: Univ. Erlangen-Nuernberg, Erlangen-Nuernberg, Fed.  
Rep. Ger.  
SOURCE: Pharmazie (1968), 23(3), 135-45  
DOCUMENT TYPE: Journal  
LANGUAGE: German

GI For diagram(s), see printed CA Issue.

AB The muscle-relaxing properties of **quaternized** amino alc. esters of 3,4-diphenyl- (I), 3,4-dimethyl- (II), 3,4-di(2-furyl)- (III), and 3,4-bis(5-nitro-2-furyl)thiophene-2,5-**dicarboxylic acid** (IV); phenanthreno[9,10-c]thiophene-1,3-**dicarboxylic acid** (V); and 2,3-diphenylbenzene-1,4- (VI), 3,6-diphenylbenzene-1,2- (VII), and 2,5-diphenylfuran-3,4-**dicarboxylic acid** (VIII) were determined. I (10 g.) was boiled with 300 ml.  $\text{SOCl}_2$  and worked up to give 8.1 g. I dichloride (IX), m. 123-4°. Similarly prepared were 37% II dichloride (X), m. 67-73°; III dichloride (XI), 91%, m. 90.5-1.5° (ligroine); IV dichloride, m. 92.5-95° ( $\text{C}_6\text{H}_6$ ); V dichloride (XII), 37%, m. 193-4° ( $\text{C}_6\text{H}_6$ ); and VI dichloride (XIII), 80%, m. 153.5-56° (decomposition) (ligroine). VIII (5.5 g.) was added in small portions with stirring to an ice-cold suspension of 16 g.  $\text{PCl}_5$  in 55 ml.  $\text{Et}_2\text{O}$ , stirred 30 min., and worked up to give 4.8 g. VIII dichloride (XIV), m. 120-1° (twice from ligroine). Crude II in dioxane was treated with  $\text{CH}_2\text{N}_2$  in  $\text{Et}_2\text{O}$ , kept 3 hrs., and worked up to give 36% di-Me ester, m. 171.5-2.5° (also prepared by heating X and  $\text{MeOH}$ ), which was refluxed in methanolic KOH and worked up to give pure II, decompose 324-7°. Similarly, III (at -5°), gave 90% di-Me ester, m. 129° (twice from  $\text{ACOH}$ ), which, at -5° in  $\text{Ac}_2\text{O}$ , was nitrated with  $\text{HNO}_3$  (d. 1.52), stirred 1 hr., and worked up to give IV di-Me ester, m. 182-4°, which refluxed 2 min. in methanolic KOH and worked up gave IV, m. 258° (decomposition). A suspension of 2 g. 1,4-dimethyl-2,3-diphenylbenzene in 60 ml.  $\text{C}_5\text{H}_5\text{N}$  and 20 ml.  $\text{H}_2\text{O}$  containing 25.3 g.  $\text{KMnO}_4$  was refluxed 2 hrs. and worked up to give 2.2 g. VI, m. 308-11°. IX (15.6 g.) and 25.2 g.  $\beta$ -diethylaminoethanol (DEAE) was refluxed 6 hrs. in 500 ml. dry  $\text{C}_6\text{H}_6$  and worked up to give 14.8 g. I bis( $\beta$ -diethylaminoethyl ester), m. 76.5-77° (ligroine); dipicrate m. 175.5-77° (1:1  $\text{Me}_2\text{CO}-\text{H}_2\text{O}$ ); di-HBr salt m. 185.5-6.5° ( $\text{Me}_2\text{CO}$ -iso- $\text{PrOH}$ ); dimethiodide m. 212-13° (decomposition); bis(benzyl bromide) decomposed 191°, m. 240-7° ( $\text{EtOH}$ - $\text{EtOAc}$ ). The following were prepared II bis( $\beta$ -diethylaminoethyl ester), 70% [di-HBr salt m. 212.5-14° (decomposition); dimethiodide m. 202.5-2.5° (decomposition)]; III bis( $\beta$ -diethylaminoethyl ester), 61%, n<sub>22D</sub> 1.459, by shaking XI and DEAE in  $\text{C}_6\text{H}_6$  66 hrs. at room temperature and working

up

[di-HBr salt, m. 179.5-81° (decomposition); dimethiodide m. 177.5-79° (decomposition)]; IV bis( $\beta$ -diethylaminoethyl ester), 47%, m. 42-7° (dimethiodide m. 192-5°); VI bis( $\beta$ -diethylaminoethyl ester), 62%, n<sub>22D</sub> 1.540 [di-HBr salt m. 185.5-7.5° ( $\text{EtOAc}$ - $\text{EtOH}$ )]; dimethiodide, m. 234-5° (decomposition)]; VIII bis( $\beta$ -diethylaminoethyl ester), 74% [di-HBr salt m. 180-1° (3:1  $\text{Me}_2\text{CO}$ - $\text{EtOH}$ )]; dimethiodide m. 185.5-87° (decomposition)] I bis( $\beta$ -dimethylaminoethyl ester) 65%, m. 69-79° (ligroine) [dimethiodide decomposed 225-50° ( $\text{EtOH}$ )]; and V bis( $\beta$ -diethylaminoethyl ester), 90%, [di-HCl salt, decompose 211-12.5°; dimethiodide m. 215-16° (decomposition)]. XII (1.25 g.) and 1.1 g.  $\text{MeOH}$  refluxed in 5 ml.  $\text{C}_6\text{H}_6$  and cooled precipitated 0.85 g. V di-Me ester, m. 118-19°. DEAE (4.7 g.) in 50 ml.  $\text{Me}_2\text{CO}$  was added to 12 g. VII anhydride suspended in 250

ml. dry refluxing Me<sub>2</sub>CO, and the mixture refluxed 20 min. to precipitate 14.6 g. of

the half **ester**, m. 205-22°, difficulty soluble in 2N NaOH and 2N HCl. This intermediate (8.35 g.) and 5.4 g.  $\beta$ -diethylaminoethyl chloride was refluxed 6.5 hrs. in 160 ml. dry iso-PrOH and worked up to give 7.3 g. VII bis( $\beta$ -diethylaminoethyl **ester**), m. 99-100° (ligroine and petroleum ether); di-HBr salt m. 193-5°; dimethiodide m. 206.5-7.5° (decomposition). The anhydride (5 g.) of *cis, cis, cis, cis*-3,6-diphenyl-1,2,3,6-tetrahydrobenzene-1,2-dicarboxylic acid in 80 ml.

HCONMe<sub>2</sub> was hydrogenated at atmospheric pressure and room temperature over Pd(OH)<sub>2</sub> on

BaSO<sub>4</sub> and worked up to give 3.3 g. anhydride of *cis, cis, cis, cis*-3,6-diphenylcyclohexane-1,2-dicarboxylic acid, m. 220-2° (EtOAc). A solution of 4.8 g. 2,7-diaminodiphenylene sulfone and 14 g. di-Et diacetylsuccinate in 20 ml. AcOH was refluxed 45 min. and cooled to precipitate 12.5 g. XV, m. 251-3° (BuOH:AcOH), saponified to the free acid by methanolic KOH. A mixture of 1.28 g. 2,2'-dihydroxy-5,5'-dimethyldeoxybenzoin in 2N NaOH and 1 g. ClCH<sub>2</sub>CO<sub>2</sub>H solution neutralized with K<sub>2</sub>CO<sub>3</sub> was refluxed 3 hrs. and worked up to give 2-hydroxy-2'-carboxymethoxy-5,5'-dimethyldeoxybenzoin, m. 159-61° (60% EtOH), and 2,2'-dicarboxymethoxy-5,5'-dimethyldeoxybenzoin, m. 172-4° (60% AcOH and 60% EtOH). Extensive biol. data are given.

IT 19799-17-8P 19799-21-4P 19799-34-9P

19799-36-1P 19802-94-9P 19802-96-1P

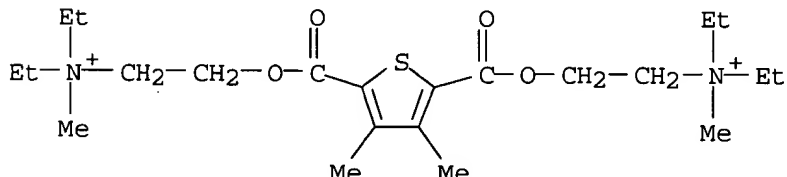
19971-00-7P 19976-53-5P 19976-55-7P

20653-69-4P

RL: SPN (Synthetic preparation); PREP (Preparation)  
(preparation of)

RN 19799-17-8 CAPLUS

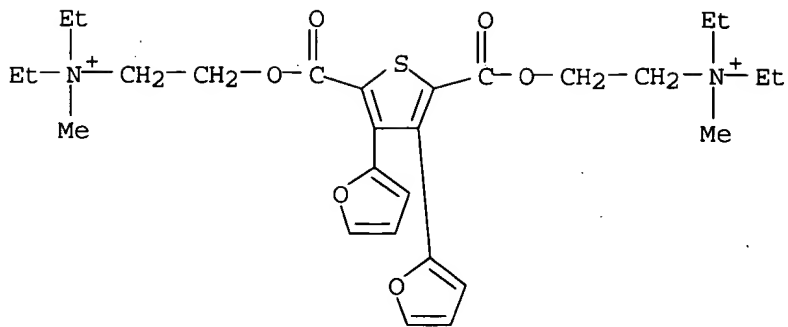
CN Ammonium, diethyl(2-hydroxyethyl)methyl-, iodide, 3,4-dimethyl-2,5-thiophenecarboxylate (2:1) (8CI) (CA INDEX NAME)



●2 I-

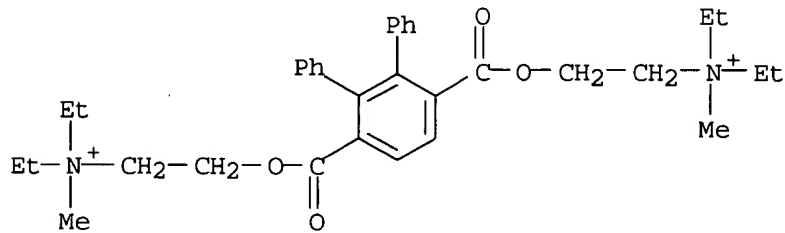
RN 19799-21-4 CAPLUS

CN Ammonium, diethyl(2-hydroxyethyl)methyl-, iodide, 3,4-di-2-furyl-2,5-thiophenedicarboxylate (2:1) (8CI) (CA INDEX NAME)



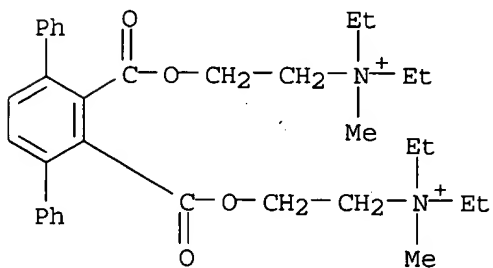
●2 I-

RN 19799-34-9 CAPLUS  
 CN Ammonium, diethyl(2-hydroxyethyl)methyl-, iodide, [o-terphenyl]-3',6'-dicarboxylate (2:1) (8CI) (CA INDEX NAME)



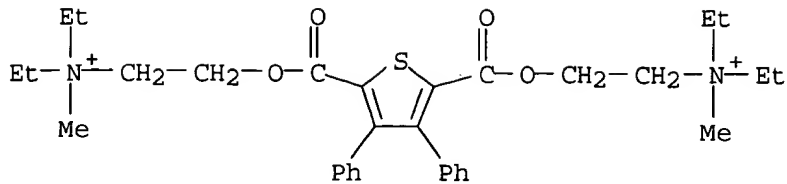
●2 I-

RN 19799-36-1 CAPLUS  
 CN Ammonium, diethyl(2-hydroxyethyl)methyl-, iodide, [p-terphenyl]-2',3'-dicarboxylate (2:1) (8CI) (CA INDEX NAME)



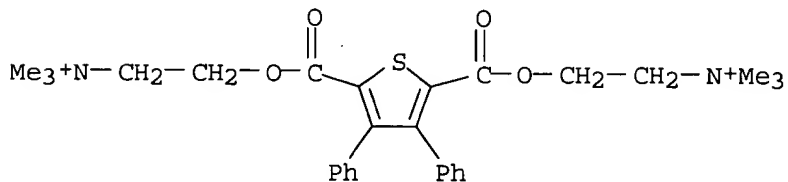
●2 I-

RN 19802-94-9 CAPLUS  
 CN Ammonium, diethyl(2-hydroxyethyl)methyl-, iodide, 3,4-diphenyl-2,5-thiophenedicarboxylate (2:1) (8CI) (CA INDEX NAME)



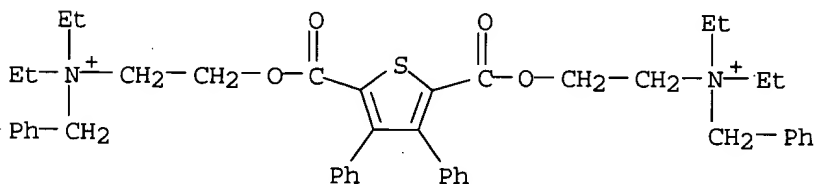
● 2 I<sup>-</sup>

RN 19802-96-1 CAPLUS  
 CN Choline, iodide, 3,4-diphenyl-2,5-thiophenedicarboxylate (2:1) (8CI) (CA INDEX NAME)



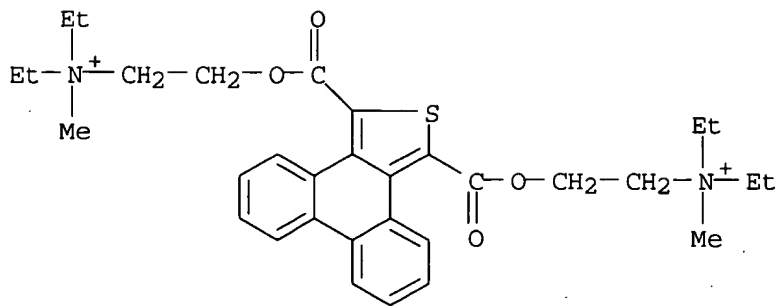
● 2 I<sup>-</sup>

RN 19971-00-7 CAPLUS  
 CN Ammonium, benzyltriethylammonium, 3,4-diphenyl-2,5-thiophenedicarboxylate (2:1) (8CI) (CA INDEX NAME)



● 2 Br<sup>-</sup>

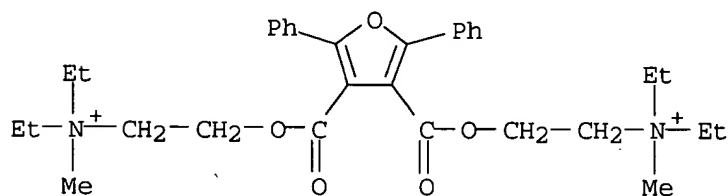
RN 19976-53-5 CAPLUS  
 CN Ammonium, diethyl(2-hydroxyethyl)methyl-, iodide, phenanthro[9,10-c]thiophene-1,3-dicarboxylate (2:1) (8CI) (CA INDEX NAME)



●2 I<sup>-</sup>

RN 19976-55-7 CAPLUS

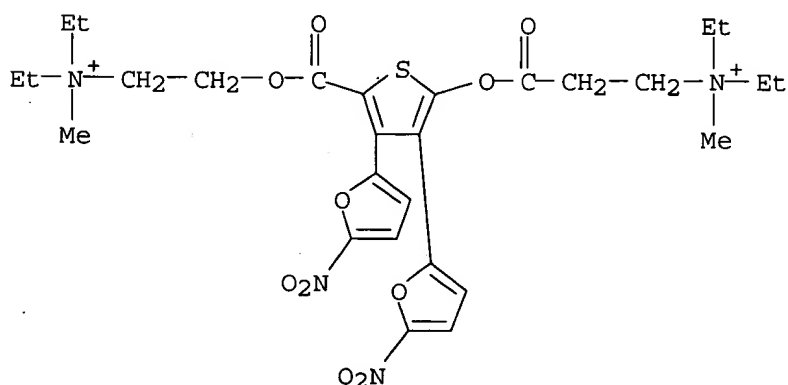
CN Ammonium, diethyl(2-hydroxyethyl)methyl-, iodide, 2,5-diphenyl-3,4-furandicarboxylate (2:1) (8CI) (CA INDEX NAME)



●2 I<sup>-</sup>

RN 20653-69-4 CAPLUS

CN Ammonium, diethyl(2-hydroxyethyl)methyl-, iodide, 3,4-bis(5-nitro-2-furyl)-2,5-thiophenedicarboxylate (2:1) (8CI) (CA INDEX NAME)



●2 I<sup>-</sup>

DOCUMENT NUMBER: 69:1562  
TITLE: Cyclobutanedicarboxylic acids. VI. Relation between curariform activity and structure in a series of alkamine cyclobutanedicarboxylic acid derivatives  
AUTHOR(S): Kharkevih, D. A.; Arendaruk, A. P.; Skoldinov, A. P.  
CORPORATE SOURCE: Mosk. Med. Inst. im. Sechenova, Moscow, USSR  
SOURCE: Khimiko-Farmatsevticheskii Zhurnal (1968), 2(3), 7-11  
CODEN: KHFZAN; ISSN: 0023-1134

DOCUMENT TYPE: Journal  
LANGUAGE: Russian

AB The curariform activity of a series of alkamine cyclobutane dicarboxylic acid derivs. was determined by drooping head symptom in rabbits and by a study of the effect of the compds. on the transfer of stimulation from the sciatic nerve to the gastrocnemius muscle of cats. Activity was studied in relation to 4 structural features: distance between the **quaternary** N atoms, radicals shielding these N atoms, stereoconfiguration of the truxilllic acids, and the structure of the aliphatic part of the mol. separating the 2 **quaternary** N atoms. The bis(N-methylpiperidino)cyclobutanedicarboxylates and bis(diethylmethylammonium)cyclobutanedicarboxylates with 11 C and 2 O atoms between the 2 **quaternary** groups were the most effective curariform agents. The  $\alpha$ -truxilllic acid derivs. were the most effective and the  $\gamma$ -truxilllic acid derivs. the least effective in suppressing transfer of nerve impulses. Replacement of **ester** groups with amides increased the curariform activity of the diphenylcyclobutanedicarboxylic acid bisquaternary ammonium salts. These compds. apparently are antidepolarizing curariform substances which interact with only 1 choline receptor. 21 references.

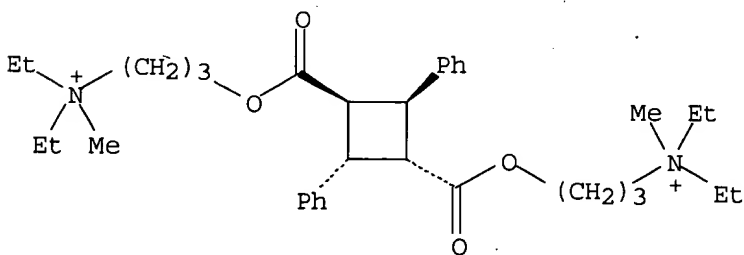
IT 4304-01-2

RL: BIOL (Biological study)  
(neuromuscular transmission inhibition by)

RN 4304-01-2 CAPLUS

CN 1-Propanaminium, 3,3'--[[(1 $\alpha$ ,2 $\alpha$ ,3 $\beta$ ,4 $\beta$ )-2,4-diphenyl-1,3-cyclobutanediyl]bis(carbonyloxy)]bis[N,N-diethyl-N-methyl-, diiodide (9CI) (CA INDEX NAME)

Relative stereochemistry.

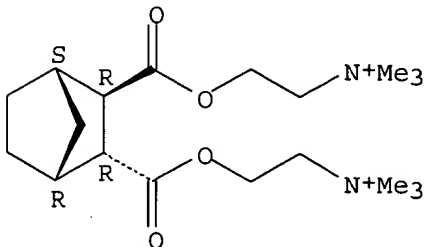


●2 I-

L9 ANSWER 7 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN  
ACCESSION NUMBER: 1966:465269 CAPLUS  
DOCUMENT NUMBER: 65:65269  
ORIGINAL REFERENCE NO.: 65:12122d-f  
TITLE: Bischoline esters of bicyclic dicarboxylic acids and related compounds  
AUTHOR(S): Koch, H.; Kotlan, J.  
CORPORATE SOURCE: Univ., Vienna

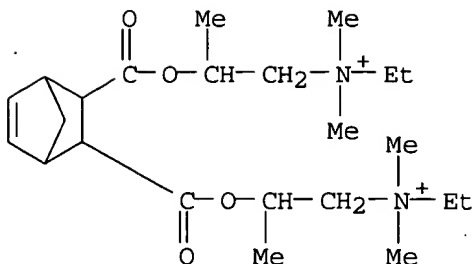
SOURCE: Monatshefte fuer Chemie (1965), 96(6),  
 2000-4  
 CODEN: MOCMB7; ISSN: 0026-9247  
 DOCUMENT TYPE: Journal  
 LANGUAGE: German  
 GI For diagram(s), see printed CA Issue.  
 AB The Me **ester** of bicyclo[2.2.1]heptane- and bicyclo[2.2.2]octane-  
 trans-dicarboxylic acids and their unsatd. analogs (I-IV) were prepared by  
 Diels-Alder reaction. The esters and dialkylamino alcs. gave basic esters  
 A (R= CHR<sub>2</sub>CH<sub>2</sub>R<sub>1</sub>), which were converted to dihydrochlorides B (R =  
 CHR<sub>2</sub>CH<sub>2</sub>R<sub>1</sub>.HCl) and bis(**quaternary** ammonium salts) C (R =  
 CHR<sub>2</sub>CH<sub>2</sub>R<sub>1</sub>.R<sub>3</sub>X) in the usual manner. The compds. prepared are given in the  
 table. Some of the compds. are muscle-relaxants. C (m.p.); R<sub>1</sub>, R<sub>2</sub>, A  
 (b.p./mm.), B (m.p.), a, b; I, NMe<sub>2</sub>, H, 180-90°/2, 203-5°,  
 234-6°, 197-9°; I, NMe<sub>2</sub>, Me, 185-95°/5,  
 215-17°, 219-21°, oil, I; NEt<sub>2</sub>, H, 205-15°/5,  
 133-5°, oil, 173-5°; I piperidino, H, 205-20°/5,  
 227-9°, 184-7°, 96-9°; II NMe<sub>2</sub>, H, 185-95°/2,  
 179-82°, 235-8°, 194-6°; II NEt<sub>2</sub>, H, 210-15°/2,  
 165-8°, -, 204-9°; III NMe<sub>2</sub>, H, 180-90°/2,  
 198-201°, 210-13°, 174-7°; III NMe<sub>2</sub>, Me,  
 190-200°/5, 231-3°, 179-82°, oil; III NEt<sub>2</sub>, H,  
 210-20°/5, 158-60°, oil, -; IV, NMe<sub>2</sub>, H, 185-90°/2,  
 175-80°, 208-11°, 217-21°; a:, R<sub>3</sub>X, =, MeI, b:, R<sub>3</sub>X,  
 =, EtBr.;  
 IT 5783-18-6, Choline, iodide, 2,3-norbornanedicarboxylate, trans-  
 7172-48-7, Ammonium, ethyl(2-hydroxypropyl)dimethyl, bromide,  
 5-norbornene-2,3-dicarboxylate, trans- 10491-44-8, Choline,  
 iodide, bicyclo[2.2.2]octane-2,3-dicarboxylate, trans-  
 (preparation of)  
 RN 5783-18-6 CAPLUS  
 CN Ethanaminium, 2,2'-[bicyclo[2.2.1]heptane-2,3-  
 diylbis(carbonyloxy)]bis[N,N,N-trimethyl-, diiodide, (2-endo,3-exo)- (9CI)  
 (CA INDEX NAME)

Relative stereochemistry.



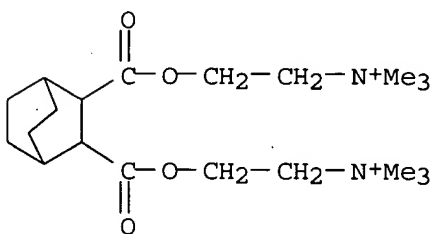
●2 I<sup>-</sup>

RN 7172-48-7 CAPLUS  
 CN Ammonium, ethyl(2-hydroxypropyl)dimethyl-, bromide, 5-norbornene-2,3-  
 dicarboxylate, trans- (8CI) (CA INDEX NAME)



● 2 Br<sup>-</sup>

RN 10491-44-8 CAPLUS  
 CN Choline, iodide, bicyclo[2.2.2]octane-2,3-dicarboxylate, trans- (8CI) (CA  
 INDEX NAME)



● 2 I<sup>-</sup>

L9 ANSWER 8 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

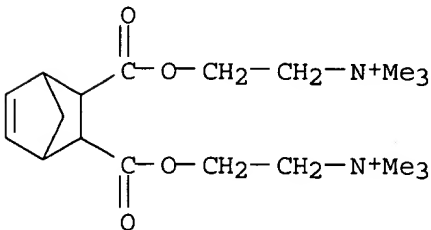
ACCESSION NUMBER: 1966:103735 CAPLUS  
 DOCUMENT NUMBER: 64:103735  
 ORIGINAL REFERENCE NO.: 64:19448h, 19449a-e  
 TITLE: Basic esters of bicyclic diacids  
 PATENT ASSIGNEE(S): Firma F. Joh. Kwizda, Heinrich Koch, Johannes Kotlan  
 SOURCE: 7 pp.  
 DOCUMENT TYPE: Patent  
 LANGUAGE: Unavailable  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
AT 244929	AT	19660210	AT	19631211 <--

AB Bicyclic diacids (I) are esterified by basic alcs. to give II. The corresponding hydrogenated esters IV and the **quaternary** salts (III.R5X) (III) and (II.R5X) (V) of II and IV resp. are prepared. All these compds. have pharmacol. properties, such as action on muscle nerves or blocking effect on ganglions. The esters are prepared from the dichlorides of I; e.g., a solution of 20 g. of dichloride of 1,4-endomethylene-2,3-trans-dicarboxy-5-cyclohexene (Ia) in 100 ml. benzene is added to 40 g. Me<sub>2</sub>NCH<sub>2</sub>CH<sub>2</sub>OH in 100 ml. benzene. The solution is refluxed, washed with 20% NaOH, and with water. Distillation gives II (n = 1, R<sub>1</sub> = R<sub>2</sub> = H, R<sub>4</sub> = R<sub>3</sub> = Me) in 95% yield, b<sub>2</sub> 180-90; n<sub>20D</sub> 1.4732; d<sub>254</sub> 1.0422. Corresponding III (n = 1) prepared are (R5X and m.p. given): HCl, 203-5°; MeI, 227-33°; EtI, 225-9°; EtBr, 191-5°. Hydrogenation of

the same II in AcOH over Pd-C gives IV (n = 1) in 80% yield: b2 175-80°; n25D 1.4729; d204 1.011. Corresponding V(n = 1) prepared are (R5X and m.p. given): HCl, 188-92°; MeI, 169-73°. Other aminoalcs. are used to give the following derivs.: II (n = 1, R3 = R4 = Et, R1 = R2 = H), b5 205-15°; n25D 1.4695; d204 0.9981; III (n = 1, R3 = R4 = Et, R1 = R2 = H (R5X and m.p. given): HCl, 149-53°; EtI, 247-53°; II (n = 1, R1 = H, R2 = R3 = R4 = Me), b5 190-5, n25D 1.4677, d204 1.0064; III (n = 1, R1 = H, R2 = R3 = R4 = Me), HCl, 228-32°; II [n = 1, R1 = R2 = H, (NR3R4) = piperidino], b5 205-25; n25D 1.4952; III [n = 1, R1 = R2 = H, NR3R4 = piperidino], HCl, deliquescent crystals; MeI, oil.. In the 1,4-endoethylene-2,3-trans-dicarboxy-5-cyclohexene series (n = 2), the following compds. are prepared by the same method (R5X and mp. given): II (n = 2, R1 = R2 = H, R3 = R4 = Me), b2 175-95; III (n = 2, R1 = R2 = H, R3 = R4 = Me, R5X = MeI), m. 235-8°; II (n = 2, R1 = R2 = H, R3 = R4 = Et), b2 210-25; III (n = 2, R1 = R2 = H, R3 = R4 = Me), HCl, 163-8°; EtBr, 204-9°; II (n = 2, R3 = R4 = Me, R4 = H, R2 = Me), b2 200-10°; III (n = 2, R2 = R3 = R4 = Me, R1 = H), HCl, 170-80°; MeI, 215-30°. Transesterification is also used to prepare II. E.g., 20 g. of the di Me ester of 5,6:7,8-dibenzo-2,3-trans-dicarboxybicyclo[2.2.2]octane (Diels-Alder adduct between di-Me fumarate and anthracene) is added to a solution of 0.5 g. Na in 40 g. Me2NCH2CH2OH and heated on a water bath 5 hrs. The more volatile fraction is then removed by vacuum, and the residue dissolved in benzene and washed with water. After drying over Na2SO4 and evaporation of the solvent, the oily ester is purified by conversion into the corresponding dihydrochloride. Dimethiodide III (n = 1, R5X = HI) (IIIA) is prepared by another method: 22 g. of the dichloride of Ia are added to an excess of cold ethylene chlorhydrin. The rough dichloroethyl ester of Ia obtained is refluxed in 300 ml. acetone with 30 g. NaI. After elimination of NaCl and removal of the solvent, the bis(iodoethyl) ester of Ia is dissolved in ether and washed with water and thiosulfate solution. The product is then heated in benzene solution with Me3N in a pressure bottle at 100° 6 hrs. IIIA is recrystd. from acetone-iso-PrOH.

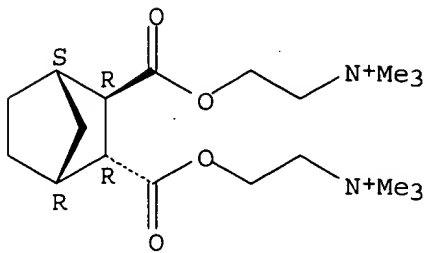
IT 5561-81-9, Choline, chloride, 5-norbornene-2,3-dicarboxylate, cis-  
 5783-18-6, Choline, iodide, 2,3-norbornanedicarboxylate, trans-  
 6012-28-8, 9,10-Ethanoanthracene-11,12-dicarboxylic  
 acid, 9,10-dihydro-, diester with choline iodide, trans-  
 (preparation of)  
 RN 5561-81-9 CAPLUS  
 CN Choline, chloride, 5-norbornene-2,3-dicarboxylate, cis- (8CI) (CA INDEX  
 NAME)



●2 Cl-

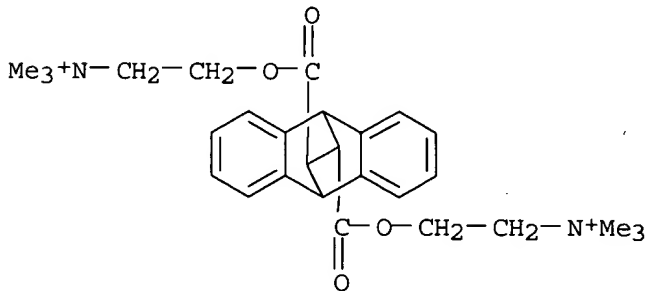
RN 5783-18-6 CAPLUS  
 CN Ethanaminium, 2,2'-[bicyclo[2.2.1]heptane-2,3-  
 diylbis(carbonyloxy)]bis[N,N,N-trimethyl-, diiodide, (2-endo,3-exo)- (9CI)  
 (CA INDEX NAME)

Relative stereochemistry.



●2 I-

RN 6012-28-8 CAPLUS  
CN Ethanaminium, 2,2'-[ (9,10-dihydro-9,10-ethanoanthracene-11,12-diyi)bis(carbonyloxy)bis[N,N,N-trimethyl-, diiodide, trans- (9CI) (CA INDEX NAME)



●2 I-

L9 ANSWER 9 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1965:455342 CAPLUS  
DOCUMENT NUMBER: 63:55342  
ORIGINAL REFERENCE NO.: 63:10134a-c  
TITLE: Plasticizer compositions  
INVENTOR(S): Kay, Ronald W.  
PATENT ASSIGNEE(S): Distillers Co. Ltd.  
SOURCE: 5 pp.  
DOCUMENT TYPE: Patent  
LANGUAGE: Unavailable  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FR 1391727	---	19650312	FR	<--
GB 1013605	---		GB	
NL 298427	---		NL	

PRIORITY APPLN. INFO.: GB 19620929

AB Alkali metal salts of an aromatic or aliphatic **dicarboxylic acid** monoester are heated with a mixture of 1,4-dichloro-2-butene (I) and a neutral alkali metal salt of an aromatic or aliphatic

**dicarboxylic acid** to give materials which can be used as plasticizers for poly(vinyl chloride) (II). Thus, 1.375 mole phthalic anhydride is dissolved in 1.52 mole BuOH at <105°, the solution is added in 1 hr. and 20 min. to a mixture of 1.25 mole I, 0.625 mole 1,4-C<sub>6</sub>H<sub>4</sub>(CO<sub>2</sub>Na)<sub>2</sub>, 0.757 mole Na<sub>2</sub>CO<sub>3</sub>, 0.016 mole Me<sub>3</sub>(PhCH<sub>2</sub>)NCl, and 312 ml. BuOH as the temperature rises from 116° to 123°, and the H<sub>2</sub>O-BuOH azeotrope is distilled. The mixture is refluxed 10 hrs., washed with H<sub>2</sub>O, washed with NaOH, washed with H<sub>2</sub>O, and distilled to give 436 g. **ester** (III) saponification number 445. III (50 parts) is incorporated in 100 parts

II to give a product, tensile strength 262.5 kg./cm.<sup>2</sup>, elongation at break 300%, melt index 19, volatilization loss 0.6%, as compared with 259, 280, 23, and 0.7, resp., for the control.

IT 3388-64-5, Ammonium, 2-butene-1,4-diylbis[triethyl-, chloride (as catalyst in esterification of 1,4-dichloro-2-butene with monobutyl esters of phthalic or succinic acids alone or with phthalic or succinic acid alkali metal salts)

RN 3388-64-5 CAPLUS

CN Ammonium, 2-butene-1,4-diylbis[triethyl-, dichloride (8CI) (CA INDEX NAME)



●2 Cl<sup>-</sup>

L9 ANSWER 10 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1964:414046 CAPLUS

DOCUMENT NUMBER: 61:14046

ORIGINAL REFERENCE NO.: 61:2368g-h,2369a-b

TITLE: Chemical and pharmacological investigations in the series of cyclobutane **dicarboxylic acid** derivatives: curarelike activity of bisquaternary salts of basic esters or amides of  $\alpha$ -,  $\epsilon$ -, and  $\gamma$ -truxillic acids

AUTHOR(S): Arendaruk, A. P.; Kravchuk, L. A.; Skoldinov, A. P.; Kharkevich, D. A.

SOURCE: Uch. Zap., Inst. Farmakol. i Khimioterapii, Akad. Med. Nauk SSSR (1963), 3, 138-57

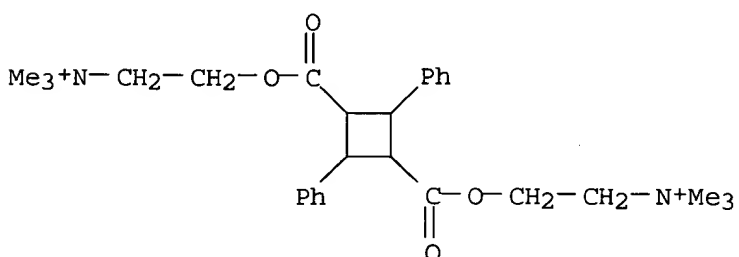
DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

AB The curarelike activities of thesin (di-d-isoretronecanol **ester** of p,p'-dihydroxy- $\alpha$ -truxillic acid; CA 54, 24835cf) and its diiodomethylate (CA 55, 15366a) were the starting point for a closer investigation of structure-activity relations of a large number of bisquaternary salts of basic esters and amides of  $\epsilon$ -(I),  $\gamma$ -(II), and  $\alpha$ -truxillic (III) acids with the scope of elucidating the influence of (1) distance between cationic centers, (2) nature of substituents at the **quaternary** N atoms, and (3) structure of the chain between cationic centers. Determination of medium effective doses (E.D.50) was carried out on rabbits by the head-drop method; blocking of excitation transmission from the sciatic nerve to the gastrocnemius muscle was studied in decerebrate cats. The [XR(CH<sub>2</sub>)<sub>n</sub>]<sub>2</sub> diiodide derivs. of III possessed greatest activity, where X = Et<sub>2</sub>N, piperidino, or 1-pyrrolidinyl, R = Me, and n = 3 and 4 (E.D.50, 25-33  $\gamma$ /kg.; transmission blocking 100-150  $\gamma$ /kg.). Compds. with n = 2, 5, 6, 7, or XMe<sub>2</sub>N, morpholino and R = Me, Et were less active, dimethiodides of dimethylaminoalkyl esters of III least active. The

influence of chain structure was studied in the dialkylaminopropylamides of I, II, and III, which possessed longer activity than the bisquaternary salts of the corresponding esters. Here also, activity decreased in the order III, I, and II derivs. All compds. investigated are nondepolarizing muscle relaxants. The X = piperidino, R = Et, n = 4 derivative, with E.D.50 = 41  $\gamma$ /kg. was proposed for clin. investigation as truxillone.

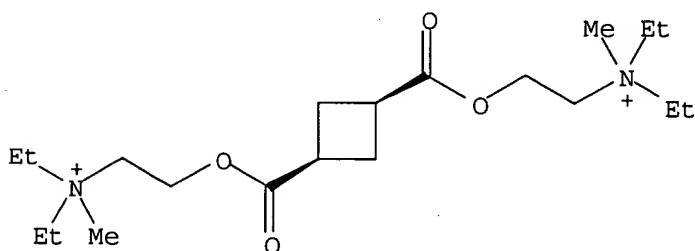
IT 10066-71-4, Choline, iodide, 2,4-diphenyl-1,3-cyclobutanedicarboxylate 17924-61-7, 1,3-Cyclobutanedicarboxylic acid, diester with diethyl(2-hydroxyethyl)methylammonium iodide, cis- (preparation, chemistry and pharmacology of)  
RN 10066-71-4 CAPLUS  
CN Choline, iodide, 2,4-diphenyl-1,3-cyclobutanedicarboxylate (2:1), cis-1,2,trans-1,3,trans-1,4- (8CI) (CA INDEX NAME)



•2 I-

RN 17924-61-7 CAPIUS  
CN Ammonium, diethyl(2-hydroxyethyl)methyl-, iodide, 1,3-cyclobutanedicarboxylate (2:1), cis- (8CI) (CA INDEX NAME)

### Relative stereochemistry.



• 2 I -

L9 ANSWER 11 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN  
ACCESSION NUMBER: 1961:124971 CAPLUS  
DOCUMENT NUMBER: 55:124971  
ORIGINAL REFERENCE NO.: 55:23573a-i,23574a-b  
TITLE: Muscarine. XI. Synthesis of bisquaternary compounds related to muscarine  
AUTHOR(S): Kiss, J.; Furter, H.; Lohse, F.; Hardegger, E.  
CORPORATE SOURCE: Eidg. Tech. Hochschule, Zurich, Switz.  
SOURCE: Helvetica Chimica Acta (1961), 44, 141-7  
CODEN: HCACAV; ISSN: 0018-019X  
DOCUMENT TYPE: Journal

LANGUAGE:

German

OTHER SOURCE(S):

CASREACT 55:124971

AB cf. CA 54, 601b. Five compds. related to muscarine were prepared from D-glucosamine-HCl (I) or L-glucosaminic acid (II). To a vibrated solution of 6 g. D-glucosaminic acid (III) in 36 ml. N HCl was added slowly during 3 hrs. at 0-5° 2.5 g. NaNO<sub>2</sub> in 40 ml. H<sub>2</sub>O. After standing 20 hrs. at 20°, the solution was evaporated in vacuo at 40°, the alc. solution of the residue was filtered, evaporated, the residue in H<sub>2</sub>O neutralized with excess CaCO<sub>3</sub>, the solution again filtered and evaporated, the viscous oil was digested six times with 40 ml. Me<sub>2</sub>CO, the residual Ca chitarate dissolved in 150 ml. H<sub>2</sub>O, and by ion exchange (Nalcite HCR) converted into chitaric acid, m. 146°, contaminated by the lactone. This acid was dissolved in a 10-fold quantity of MeOH and treated 24 hrs. with excess Me<sub>2</sub>NH at room temperature. Evaporation in vacuo gave chitaric acid dimethylamide

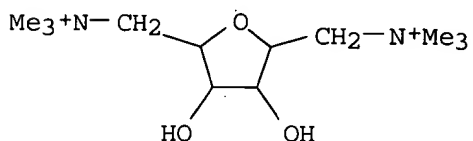
(IV), m. 172° (EtOAc), [α]D 18.8° (c 1, MeOH). IV (2.4 g.) in 11 ml. HNO<sub>3</sub> (d. 1.2) was carefully warmed to 65°. A vigorous reaction occurred, ending after 30-5 min. After repeated evapns. of the aqueous solns., the remaining viscous oil was crystallized from Me<sub>2</sub>CO to yield 63% 2,5-anhydro-D-saccharic acid 1-dimethylamide (V), decomposing at 203-4°, [α]D -2.4° (c 2.5, H<sub>2</sub>O). From the mother liquor 2,5-anhydro-D-saccharic acid (VI) was isolated via the Ca salt (yield 0.32 g.), m. 170-1° (Me<sub>2</sub>CO-Et<sub>2</sub>O), [α]D 39.3° (c 2, H<sub>2</sub>O). Sublimation of V in high vacuum at 220° gave 2,5-furandicarboxylic acid (VII), decomposing at 300-20°. Oxidation of 15 g. I with 41 ml. HNO<sub>2</sub> (d. 1.2) at 60° gave via the Ca salt 19% crude 2,5-anhydro-D-manno-saccharic acid (VIII), m. 182° (alc.-Et<sub>2</sub>O), [α]D 46.3° (c 1, H<sub>2</sub>O). Esterification of VIII in MeOH with ethereal CH<sub>2</sub>N<sub>2</sub> yielded quant. the di-Me **ester** hemihydrate (IX), m. 71° (MeOH-Et<sub>2</sub>O), [α]D 45.3° (c 2.5, H<sub>2</sub>O). IX (3.1 g.) in 10 ml. MeOH with 5 g. Me<sub>2</sub>NH 2 hrs. at 65° in an autoclave yielded 2.2 g. VIII bis(dimethylamide) (X), m. 126° (EtOAc), [α]D 93.3° (c 1.5, H<sub>2</sub>O). Esterification of V in MeOH with excess ethereal CH<sub>2</sub>N<sub>2</sub> yielded V 6-Me **ester** (XI), m. 165-6° (EtOAc), [α]D 11.5° (c 1, H<sub>2</sub>O). XI (0.4 g.) in 5 ml. cold MeOH treated with 5 g. Me<sub>2</sub>NH in the cold and kept 24 hrs. at room temperature yielded 0.34 g. 2,5-anhydro-D-saccharic acid bis(dimethylamide) (XII), decomposing at 202° (MeOH), [α]D -52.2° (c 2, H<sub>2</sub>O). XII was also prepared from 0.3 g. VI via the di-Me **ester**; yield 0.28 g. From 1.2 g. XI in 10 ml. MeOH, saturated with NH<sub>3</sub> at 0°, 0.91 g. 2,5-anhydro-D-saccharic acid 1-dimethylamide 6-**amide** was obtained, decomposing at 234°, [α]D 3.6° (c 3.5, H<sub>2</sub>O). Treatment of 2.5 g. XII in 15 ml. dry pyridine with 0.9 g. tosyl chloride in 15 ml. pyridine 24 hrs. at room temperature and after warming to 35° addition of 40 ml. EtOAc yielded 170 mg. 3,4-ditosyl-2,5-anhydro-D-saccharic acid 1,6-bis(dimethylamide) (XIII), m. 202-3° (CHCl<sub>3</sub>-EtOAc), [α]D 7.8° (c 0.65, CHCl<sub>3</sub>). Ca salt of VII (5 g.) in 150 ml. H<sub>2</sub>O was hydrogenated 6 hrs. with 5 g. Raney Ni at 150°/135 atmospheric in an autoclave. After removal of the Ca<sup>++</sup> ions by ion exchange (Nalcite HCR), 55% cis-tetrahydrofuran-2,5-dicarboxylic acid (XIV) was obtained, m. 125-6° (EtOAc-petr. ether). XIV was converted into its di-Me **ester** (XV) by the diazomethane method. Treatment of 1.02 g. XV in 10 ml. MeOH with 2 g. Me<sub>2</sub>NH 2 hrs. at 90° gave 920 mg. XIV bis(dimethylamide) (XVI). The bis(dimethylamides) X, XII, XIII, and XVI were reduced by slow addition of these compds. in dioxane to LiAlH<sub>4</sub> in dioxane, the excess LiAlH<sub>4</sub> decomposed by EtOAc, the solution evaporated, 50 ml. 10N NaOH/g. LiAlH<sub>4</sub> added,

the

tertiary amines repeatedly extracted with Et<sub>2</sub>O, and after evaporation **quaternized** by boiling in excess MeI. Thus, 1.5 g. X in 80 ml. dioxane gave after 3 hrs. reflux, etc., 1.26 g. 1,6-dideoxy-1,6-bis(dimethylamino)-2,5-anhydro-D-mannitol-2MeI, decomposing at 299-300° (80% alc.), [α]D 32.2° (c 1, H<sub>2</sub>O). XII (0.5

g.) gave 510 mg. 1,6-dideoxy-1,6-bis(dimethylamino)-2,5-anhydro-D-sorbitol-2MeI (XVII), decomposing at 300° (75% alc.),  $[\alpha]_D$  13.6° (c 1, H<sub>2</sub>O). Reduction of XIII gave two products because of partial or total reductive cleavage of the tosyl groups. Thus, 114 mg. XIII in 30 ml. dioxane with 200 mg. LiAlH<sub>4</sub> in 40 ml. dioxane gave after 2 hrs. at 80-90° and 1 hr. reflux 68 mg. crude methiodide. The fraction, difficultly soluble in alc., gave after several recrystns. from 80% alc. 3- or 4-deoxy analog of XVII, decomposing at 298-300°  $[\alpha]_D$  2.8° (c 0.7, H<sub>2</sub>O). The fraction, well soluble in alc., gave 18 mg. cis-2,5-bis(dimethylaminomethyl)tetrahydrofuran-2MeI (XVIII), decomposing at 302° (alc.), showing no rotation. The structure of XVIII was confirmed by the isolation of an identical product from the reduction of XVI in the above way. The L-form of XVII was prepared in exactly the same way as described for XVII, starting from II, with corresponding values of the phys. consts.

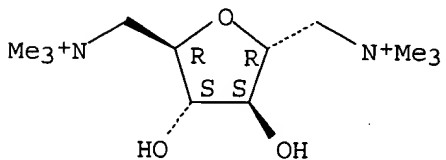
IT 88777-26-8, Sorbitol, 2,5-anhydro-1,6-dideoxy-1,6-bis(dimethylamino)-, dimethiodide 109215-30-7, Mannitol, 2,5-anhydro-1,6-dideoxy-1,6-bis(dimethylamino)-, dimethiodide (preparation of)  
 RN 88777-26-8 CAPLUS  
 CN D-Glucitol, 2,5-anhydro-1,6-dideoxy-1,6-bis(trimethylammonio)-, diiodide (9CI) (CA INDEX NAME)



●2 I<sup>-</sup>

RN 109215-30-7 CAPLUS  
 CN Mannitol, 2,5-anhydro-1,6-dideoxy-1,6-bis(dimethylamino)-, dimethiodide (6CI) (CA INDEX NAME)

Absolute stereochemistry.



●2 I<sup>-</sup>

L9 ANSWER 12 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1959:94742 CAPLUS  
 DOCUMENT NUMBER: 53:94742  
 ORIGINAL REFERENCE NO.: 53:17092f-i,17093a-g  
 TITLE: Pyrroles. XIV. Mannich bases of 2,5-substituted pyrroles  
 AUTHOR(S): Herz, Werner; Settine, Robert L.  
 CORPORATE SOURCE: Florida State Univ., Tallahassee  
 SOURCE: Journal of Organic Chemistry (1959), 24,

DOCUMENT TYPE:

Journal

LANGUAGE:

Unavailable

GI For diagram(s), see printed CA Issue.

AB cf. C.A. 53, 7136h. Several pyrroles substituted in the 2-and 5-positions of the pyrrole nucleus were subjected to the Mannich reaction. The resulting bases were shown to be 3-dialkylaminomethyl- and 3,4-bis(dialkylaminomethyl) derivs. of pyrrole. Their utility as alkylating agents was investigated. The following general procedures were used for the preparation of the Mannich bases,  $\text{Me}_2\text{NCH}_2\text{C:CR.NR}'\text{CR:CH}$  (I) and  $\text{Me}_2\text{NCH}_2\text{C:CR.NR}'\text{CR:CCH}_2\text{NMe}_2$  (II). Method A.  $\text{NHMe}_2\cdot\text{HCl}$  (85 g.) in 79 g. 40% HCHO added at  $60^\circ$  to 100 g. 2,5-dimethylpyrrole, diluted with  $\text{H}_2\text{O}$ , extracted with  $\text{Et}_2\text{O}$ , and the aqueous layer poured into 200 ml. 25% NaOH

gave

148 g. product. For disubstitution, the quantities of  $\text{NHMe}_2\cdot\text{HCl}$  and HCHO were doubled. Method B.  $\text{NHMe}_2$  (20 ml., 33%) and 20 ml. AcOH mixed with 8.5 ml. 40% HCHO, and the solution added dropwise under N to 17.1 g.

1-phenyl-2,5-dimethylpyrrole gave 11.1 g. product. For disubstitution, 2 moles aqueous  $\text{NHMe}_2$  and 2 moles HCHO were used. The picrates were precipitated by

mixing alc. solns. of the base and picric acid and recrystg. from alc. Methiodides were prepared by addition of I or II, dissolved in a min. of alc., to 10% excess MeI with stirring at ice bath temperature and recrystg. from alc. (type of compound, R, R', % yield, m.p. or b.p./mm., method, m.p. of MeI derivative, and m.p. of picrate given): I, Me, H, 92, 99-100°, A, 130° (decomposition), 117-18°; I, Me, Ph, 49.5, 130-1°/1, B, 211-12° (decomposition), 137-8°; I, Me, Me, 69, 73-4°/1, B, 140° (decomposition), 137-8° (decomposition); I, Ph, H, 73, 124-5% B, -, 179-80°; II, Me, H, 90, 144-5°, A, 139-40°, 139-40°; II, Me, Ph, 61, 150°/1, B, 100° (decomposition), 200-1° (decomposition); II, Me, Me, 72.5, 96-7°/0.3, B, -, 181-2° (decomposition). II (R = Me, R' = H) (20 g.) in 100 ml. alc. heated 48 hrs. at  $100^\circ$  with 4 g. Raney Ni and H at 80-100 atmospheric gave 7.7 g. 2,3,4,5-tetramethylpyrrole, m. 107-8°, by steam distillation. In a similar manner hydrogenolysis of 20 g. I (R = Me,

R'

= H) 8 hrs. at 80-90° gave 1.5 g. 2,3,5-trimethylpyrrole, b15 79-80°, and 7.5 g. of starting material. Di-Et acetamidomalonate (III) (16.2 g.) and 11.5 g. I (R = Me, R' = H) mixed with 100 ml. alc. containing 1.72 g. Na, treated dropwise at  $35^\circ$  with 15.8 g.  $\text{Me}_2\text{SO}_4$ , stirred overnight, and concentrated gave 18.4 g. di-Et 2,5-dimethyl-3-pyrrolylmethyl- $\alpha$ -acetamidomalonate (IV), m. 176-7°

(alc.- $\text{H}_2\text{O}$ ). Reaction of 26 g. I with 32.4 g. III in 300 ml. PhMe containing 1 g. powdered NaOH gave 24 g. IV. Condensation of 20.3 g. di-Et formamidomalonate (IVa) and 11.5 g. I (R = Me, R' = H) in alc. by quaternization in situ gave 19.2 g. di-Et 2,5-dimethyl-3-pyrrolylmethyl- $\alpha$ -formamidomalonate (V), m. 137-8.5°.

Hydrolyzing 6 g. V with 50 ml. 25% NaOH 2.5 hrs., cooling, acidifying, filtering, acidifying the filtrate to pH 5, and seeding gave 3.3 g. crude material which was chromatographed to show one spot; the analysis indicated the presence of inorg. material which could not be removed.

$\text{CH}_2(\text{CO}_2\text{Et})_2$  (VI) (50 g.) with 31.5 g. I (R = Me, R' = H) by

quaternization gave 35.5 g. di-Et 2,5-dimethyl-3-pyrrolylmethylmalonate, b2 173-5°. Reaction of 20.8 g.

2,5-diphenyl-3-(dimethylaminomethyl)pyrrole and 20.3 g. IVa by the in situ quaternization method gave 31.05 g. di-Et 2,5-diphenyl-3-pyrrolylmethyl- $\alpha$ -formamidomalonate (VII), m. 164-5°

(alc.- $\text{H}_2\text{O}$ ). VII (5 g.), 10 g. KOH, and 50 ml. 80% alc. refluxed overnight gave 1.5 g. 2,5-diphenyl-3-pyrrolealanine, m. 217-18° (decomposition).

VI (64 g.) containing 1.84 g. Na heated 6 hrs. at  $120^\circ$  under N with 26

g. I.MeI (R = Me, R' = Ph),  $\text{H}_2\text{O}$  added, and the mixture extracted with  $\text{Et}_2\text{O}$  and distilled gave 23.6 g. VI and 11.4 g. di-Et 1-phenyl-2,5-dimethyl-3-

pyrrolylmethylmalonate, b0.1 164-5°, n25D 1.5230. By the above procedure there was obtained after 24 hrs. 6.8 g. di-Et 1,2,5-trimethyl-3-pyrrolylmethylmalonate, b0.9 145-6°, n25D 1.4670. I.MeI (R = Me, R' = Me) (55 g.), 30 g. NaCN, and 200 ml. H2O heated under N until evolution of basic gas ceased gave 5.5 g. 1,2,5-trimethyl-3-pyrroleacetonitrile (VIII), b0.2 90°, n25D 1.1527,  $\nu$  2250 cm.-1. The pot residue consisted of a tarry solid which liberated NH3 on treatment with base and probably contained some amide, due to partial hydrolysis of VIII. VIII (5 g.), 5 g. KOH, and 50 ml. 80% alc. refluxed 8 hrs., diluted with H2O, and poured over ice containing 10 ml. concentrated HCl, and the oil which separated taken up in Et2O gave

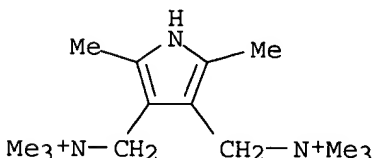
3.2 g. 1,2,5-trimethyl-3-pyrroleacetic acid (IX), m. 120-1° (ligroine). 1,2,5-Trimethylpyrrole (90 g.) and 4 g. Cu powder treated dropwise with 48 g. N2CHCO2Et, stirred 3 hrs., the Cu removed, and the filtrate distilled in vacuo gave 72.5 g. unchanged pyrrole and 18.8 g. Et 1,2,5-trimethyl-3-pyrroleacetate (X), b5 124-5°, n27D 1.4919. Hydrolysis of X with 80% alc. alkali gave 2.76 g. IX. I.MeI (R = Me, R' = Ph) (40 g.) and 30 g. NaCN similarly gave 9 g. 1-phenyl-2,5-dimethyl-3-pyrroleacetonitrile (XI), b0.6 144-5°, n25D 1.5246,  $\nu$  2250 cm.-1 (CN band). XI (3 g.) on saponification gave 2.5 g. 1-phenyl-2,5-dimethyl-3-pyrroleacetic acid, m. 151-2° (ligroine).

IT 109286-53-5, Ammonium, [(2,5-dimethylpyrrole-3,4-diyl)dimethylene]bis[trimethyl- iodide]

(preparation of)

RN 109286-53-5 CAPLUS

CN [(2,5-Dimethylpyrrole-3,4-diyl)dimethylene]bis[trimethylammonium iodide] (6CI) (CA INDEX NAME)



●2 I-

L9 ANSWER 13 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1959:94484 CAPLUS

DOCUMENT NUMBER: 53:94484

ORIGINAL REFERENCE NO.: 53:16994b-i,16995a-f

TITLE: Synthesis of some conjugated cyclobutane polyolefins and their 1,2-cycloaddition to tetracyanoethylene

AUTHOR(S): Blomquist, A. T.; Meinwald, Yvonne C.

CORPORATE SOURCE: Cornell Univ., Ithaca, NY

SOURCE: Journal of the American Chemical Society (1959), 81, 667-72

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

OTHER SOURCE(S): CASREACT 53:94484

AB 3-Methylene-1,4-diphenyl-2-methylcyclobutene (I) and diphenyldimethylenecyclobutene (II) were synthesized from  $\beta$ -truxinic acid (III) by a series of conventional transformations which included Hofmann degradation of appropriate bis(**quaternary** hydroxides) in the final steps. I and II reacted with (NC)2C:C(CN)2 (IV) by 1,2-cycloaddn. to yield spirocyclobutane derivs. 3-Methylenecyclohexene

(V) showed similar 1,2-cycloaddn. IV and norbornadiene (VI) also formed a 1:1 adduct. III (64 g.), 300 cc. absolute EtOH, 150 cc. PhMe, and 3 cc. concentrated H<sub>2</sub>SO<sub>4</sub> refluxed azeotropically and distilled gave 71.9 g. di-Et ester (VII) of III, m. 51°. VII (67.7 g.) in 100 cc. dry Et<sub>2</sub>O reduced with 14.8 g. LiAlH<sub>4</sub> in 750 cc. dry Et<sub>2</sub>O yielded 47.5 g. 1,2-bis(hydroxymethyl)-3,4-diphenylcyclobutane (VIII), m. 110-11° (MeOH). VIII (13.4 g.) added in several portions with stirring and cooling to 13.7 g. PBr<sub>3</sub>, stirred at room temperature to solution, heated 8 hrs. at

80°, cooled, diluted with 25 cc. H<sub>2</sub>O, extracted with C<sub>6</sub>H<sub>6</sub>, the extract washed, dried, evaporated, and the residue recrystd. from CCl<sub>4</sub>-pentane yielded 15.2 g. 1,2-bis(bromomethyl)-3,4-diphenylcyclobutane (IX), 95.5-6.5°. IX (3.94 g.), 4.8 g. Me<sub>3</sub>N, and 2 cc. MeOH kept 1 week at room temperature in a sealed tube gave 5 g.

1,2-bis(dimethylaminomethyl)-3,4-diphenylcyclobutane dimethobromide (X), characterized as the dipicrate, m. 255-6° (decomposition) (EtOH). X (2.56 g.) in 10 cc. H<sub>2</sub>O treated with Ag<sub>2</sub>O from 3.4 g. AgNO<sub>3</sub> and 1.4 g. KOH, stirred 2 hrs., filtered, evaporated, the residual glassy solid heated at 120-40°/0.4-0.5 mm., and the sublimate (0.75-0.9 g.) resublimed at 55°/0.4 mm. and recrystd. from MeOH gave I, needles, m. 63-4°. I in EtOAc hydrogenated at 25°/736.5 mm. over prereduced PtO<sub>2</sub> during 1 hr. and the crude product chromatographed on Al<sub>2</sub>O<sub>3</sub> gave 2,3-diphenyl-1,4-dimethylcyclobutene, n<sub>25</sub>D 1.5892. I (114.3 mg.) in 30 cc. CH<sub>2</sub>Cl<sub>2</sub> treated with ozonized O at -78° during 0.5 hr., added with stirring to 0.2 g. Zn dust in 10 cc. AcOH, stirred 0.5 hr. at room temperature, distilled into

200

mg. dimedon, a drop piperidine, and 10 cc. 75% EtOH, and the distillate heated until all CH<sub>2</sub>Cl<sub>2</sub> was removed and cooled gave 35 mg. dimedon derivative of CH<sub>2</sub>O, needles, m. 190-1°. IX (3.94 g.), 1.78 g.

N-bromosuccinimide, a few crystals of Bz<sub>2</sub>O<sub>2</sub>, and 100 cc. CCl<sub>4</sub> refluxed 1 hr. and filtered, the filtrate concentrated to about 10 cc., and diluted with pentane gave 3.4 g. 3-Br derivative (XI) of IX, plates, m. 105-5.5° (decomposition). XI and excess Me<sub>3</sub>N heated 2 days at 50° in MeOH in a sealed tube and the mixture evaporated gave 4.5 g. mixture of the 2(or

3)-butene

analog (XII) and the 3(or 2)-analog (XIII) of X; the mixture extracted with CH<sub>2</sub>Cl<sub>2</sub> left 1.5 g. of one of the isomers, m. 200-4° [picrate m. 228-30° (decomposition) (EtOH)]; the extract evaporated and the residue recrystd. from H<sub>2</sub>O yielded the other isomer, m. 185-7°, which did not yield a solid picrate. Mixture (2.04 g.) of XII and XIII converted in the usual manner to the base mixture and the crude product (1.21 g.) pyrolyzed gave 0.28-0.35 g. crude product which resublimed at 40°/0.3 mm. gave II, m. 42-3°; II could be kept several days at 0° under N without visible change but it turned yellow at room temperature within a few hrs.; in all pyrolyses 0.25-0.3 g. dark polymeric residue was formed; it could be repptd. from C<sub>6</sub>H<sub>6</sub> with hexane. II hydrogenated over PtO<sub>2</sub> absorbed 90% of 3 equivs. H. The reductive ozonolysis of II yielded 34% CH<sub>2</sub>O. II (0.32 g.) in 2 cc. CCl<sub>4</sub> titrated at 0° with 10% Br in CCl<sub>4</sub> and evaporated in vacuo gave 0.75 g.

1,2-dibromo-1,2-bis(bromomethyl)-3,4-diphenyl-3-cyclobutene, m. 118-19° (Et<sub>2</sub>O). II did not react at 25° with maleic anhydride, N-phenylmalimide, and (.tplbond.CCO<sub>2</sub>Et)<sub>2</sub> (XIV), but polymerization occurred in all cases at higher temps.; II and XIV heated at 150° and the crude product chromatographed gave a small amount of amorphous product, decompose 160-70°. II (0.34 g.), in a few cc. C<sub>6</sub>H<sub>6</sub> treated under N with 0.35 g. IV in C<sub>6</sub>H<sub>6</sub>, refluxed 0.5 hr., kept at room temperature overnight, evaporated in vacuo, the dark residue extracted with boiling

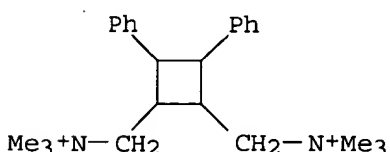
Et<sub>2</sub>O to leave an insol. polymeric residue, and the extract treated with Norite and evaporated gave 0.18 g. 1,1,2,2-tetracyano-5,6-diphenyl-7-methylenespiro[3.3]-5-heptene (XV), needles, m. 175-6° (decomposition). XV (47 mg.) in 0.5 cc. CHCl<sub>3</sub> titrated with Br in CHCl<sub>3</sub>, the solution evaporated,

and the crude residue recrystd. from Et<sub>2</sub>O gave 1,1,2,2-tetracyano-5-bromo-5-(bromomethyl)-6,7-diphenylspiro[3.3]-6-heptene, m. 162.5-63° (decomposition). I (0.2144 g.) in 5 cc. dry Et<sub>2</sub>O added to 0.120 g. IV in 10 cc. Et<sub>2</sub>O, shaken 3 hrs. at room temperature, and evaporated in vacuo gave 1,1,2,2-tetracyano-5,7-diphenyl-5-methylspiro[3.3]-5-heptene, light yellow glass, m. 139.5-40.5° (decomposition) (Et<sub>2</sub>O-petr. ether). Crude 2-cyclohexenemethanol (XVI) (13 g.) and 17.3 g. phthalic anhydride in 15 cc. PhMe refluxed 3 hrs., kept at room temperature overnight, diluted with Et<sub>2</sub>O, filtered, the filtrate extracted with aqueous Na<sub>2</sub>CO<sub>3</sub>, the alkaline extract acidified and extracted with CH<sub>2</sub>Cl<sub>2</sub>, the extract filtered and evaporated, and the residue recrystd. from hexane-Et<sub>2</sub>O gave acid phthalate (XVII) of XVI, m. 73.5-5.5° (hexane-Et<sub>2</sub>O). XVII (32 g.) in 180 cc. 25% aqueous NaOH refluxed 2 hrs. gave 12.7 g. pure XVI, b<sub>8</sub> 95°, n<sub>25</sub>D 1.4820, which treated with Ac<sub>2</sub>O and C<sub>5</sub>H<sub>5</sub>N at room temperature gave 16.7 g. acetate (XVIII) of XVI, b<sub>15</sub> 95-6°, n<sub>25</sub>D 1.4575. XVIII pyrolyzed at 525 ± 15° gave 6.9 g. crude pyrolyzate which fractionated gave V, b. 109-10°, n<sub>25</sub>D 1.4895. V (1.6 g.) added to 1.6 g. IV in C<sub>6</sub>H<sub>6</sub>, refluxed 10 min., kept at room temperature overnight, filtered, the filtrate evaporated, and the residue recrystd. from Et<sub>2</sub>O yielded 1.4 g. 1,1,2,2-tetracyanospiro[3.5]-5-nonene, m. 121-2° (decomposition) with softening and yellowing at 100-16° depending on the rate of heating. IV in C<sub>6</sub>H<sub>6</sub> treated with the usual fashion with VI and the mixture refluxed 0.5 hr. yielded 100% 8,8,9,9-tetracyanoquadracyclo[2.2.1.0.2,6.23,5]nonane (XIX), m. 186-8° (decomposition) (C<sub>6</sub>H<sub>6</sub>); when the addition was carried out at room temperature during 3-4 days a lower melting form, m. 158-60°, of XIX was obtained in 100% yield; the lower melting form changed after standing at room temperature for more than 1 week to the higher melting modification. XIX (1.3 g.) refluxed 24 hrs. with 10 g. NaOH in 12 cc. H<sub>2</sub>O and 30 cc. EtOH, acidified, concentrated, diluted with 10 cc. H<sub>2</sub>O, extracted with Et<sub>2</sub>O, and the glassy residue (1.6 g.) recrystd. from C<sub>6</sub>H<sub>6</sub>-Et<sub>2</sub>O yielded 1.25 g. 8,9-dicarboxyquadracyclo[2.2.1.0.2,6.23,5]nonane-8,9-dicarboximide, m. 205-7° (with effervescence). Methylene-1,2-cyclopropanedicarboxylic acid, methylenecyclobutane, methylenecyclononane, and norbornene added to saturated solns. of IV in C<sub>6</sub>H<sub>6</sub>, kept at room temperature overnight, and heated several hrs. at 80-90° gave only unchanged starting materials. The infrared absorption spectra of I and II are recorded.

IT 121447-16-3, Ammonium, [(3,4-diphenyl-1,2-cyclobutylene)dimethylene]bis[trimethyl-] 122568-22-3, Ammonium, [[3,4-diphenyl-3-cyclobuten-1,2-ylene]dimethylene]bis[trimethyl-] 122568-24-5, Ammonium, [[3,4-diphenyl-2-cyclobuten-1,2-ylene]dimethylene]bis[trimethyl-] (-salts)

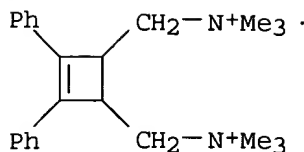
RN 121447-16-3 CAPLUS

CN 1,2-Cyclobutanedimethanaminium, N,N,N,N',N',N'-hexamethyl-3,4-diphenyl- (9CI) (CA INDEX NAME)



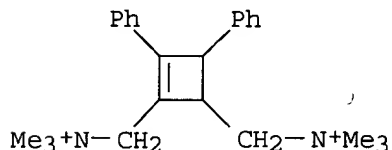
RN 122568-22-3 CAPLUS

CN 3-Cyclobutene-1,2-dimethanaminium, N,N,N,N',N',N'-hexamethyl-3,4-diphenyl- (9CI) (CA INDEX NAME)



RN 122568-24-5 CAPLUS

CN 2-Cyclobutene-1,2-dimethanaminium, N,N,N',N'',N''-hexamethyl-3,4-diphenyl- (9CI) (CA INDEX NAME)



L9 ANSWER 14 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1959:72288 CAPLUS

DOCUMENT NUMBER: 53:72288

ORIGINAL REFERENCE NO.: 53:13065b-d

TITLE: Curare-like **quaternary** salts of basic esters of aliphatic dicarboxylic acids

INVENTOR(S): Wunderlich, Helmut

DOCUMENT TYPE: Patent

LANGUAGE: Unavailable

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

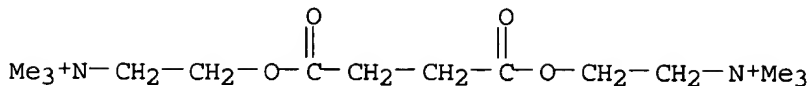
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DD 11654	-----	19560602	DD	-----

AB Bis- $\omega$ -haloalkylesters (I) of aliphatic dicarboxylic acids (II) are treated with gaseous Me3N in an inert solvent (e.g., Et2O, Me2CO, C6H6) to give **quaternary** salts. I and II are prepared from dihalides (preferably dibromides) of the acids with ethylene oxide. The **dicarboxylic acid** dichlorides may be converted into the dibromides by heating with gaseous HBr. E.g., 310 g. succinyl dichloride is brominated with HBr at 45° until the weight is 500-20 g., then fractionated to yield 85-90% succinyl dibromide III, b12 108-14°. III (488 g.) and a few crystals ZnCl2 is treated with 200 g. ethylene oxide and fractionated to give 85-90% succinylbis(bromoethyl) **ester**, b3.5 163-70° (IV). IV (166 g.) is dissolved in 400 cc. dry Me2CO and treated with a minute excess of Me3N gas yielding crystalline succinylbischoline **ester** dibromide, m. 225-7°, yield 80-5% (MeOH).

IT 55-94-7, Choline, bromide succinate  
(preparation of)

RN 55-94-7 CAPLUS

CN Ethanaminium, 2,2'-(1,4-dioxo-1,4-butanediyl)bis(oxy)bis[N,N,N-trimethyl-, dibromide (9CI) (CA INDEX NAME)



●2 Br-

L9 ANSWER 15 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1958:50718 CAPLUS

DOCUMENT NUMBER: 52:50718

ORIGINAL REFERENCE NO.: 52:9173i, 9174a-c

TITLE: Multivalent **quaternary** ammonium compounds.

VI. Some reaction products of bile acids and sterols

Lettre, H.; Gottstein, W.; Scholtissek, Ch.

CORPORATE SOURCE: Univ. Heidelberg, Germany

SOURCE: Monatshefte fuer Chemie (1957), 88, 715-20

CODEN: MOCMB7; ISSN: 0026-9247

DOCUMENT TYPE: Journal

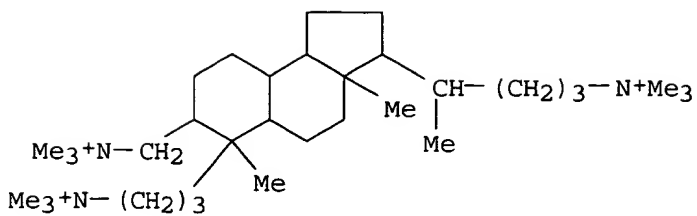
LANGUAGE: Unavailable

AB cf. C.A. 51, 4409a. Some N derivs. of lithiobilanic acid (I) and sitosterol are prepared I is treated with Ac<sub>2</sub>O followed by Me<sub>2</sub>NH to yield lithiobilanic acid 3-monodimethylamide, m. 251-2°. I with Ac<sub>2</sub>O followed by PC15 and then with Me<sub>2</sub>NH in Et<sub>2</sub>O gives an Et<sub>2</sub>O phase containing 60%-70% I. 3,4,24-tris(dimethylamide), m. 151-2°, purified by chromatography on Al<sub>2</sub>O<sub>3</sub>. The aqueous phase of the reaction yields 15-20% of I 3,24-bis(dimethylamide) (II), m. 232-3°. II is esterified with CH<sub>2</sub>N<sub>2</sub> and reduced with LiAlH<sub>4</sub> in tetrahydrofuran to 3,4-secocholan-4-ol-3,24-bis(dimethylamine hydrochloride), m. 292-5° (decomposition). II is similarly reduced to 90% 3,4-secocholane-3,4,24-tris(dimethylamine hydrochloride), decompose 275°, which forms 3,4-secocholane-3,24-tris(trimethylammonium iodide), m. 290° (decomposition). The **dicarboxylic acid** of sitosterol (III), heated 2 hrs. with Ac<sub>2</sub>O gives 76% 2,3-secositostanol-2,3-**dicarboxylic acid** anhydride, m. 176°. III di-Me **ester** is reduced by LiAlH<sub>4</sub> to 88% 2,3-secositostane-2,3-diol, m. 182-3° (MeOH). III with PC15 and Me<sub>2</sub>NH yields by chromatography on Al<sub>2</sub>O<sub>3</sub> 48% 2,3-secositostane-2,3-**dicarboxylic acid** dimethylamide, m. 106-7°, reduced by LiAlH<sub>4</sub> to 68% 2,3-secositostane-2,3-bis(dimethylamine hydrochloride), m. 326° (decomposition). This compound with MeI gives 2,3-secositostane-2,3-bis(trimethylammonium iodide), m. 323°.

IT 122387-46-6, 3,4-Secocholane-3,4,24-triamine, N<sub>3</sub>,N<sub>3</sub>,N<sub>4</sub>,N<sub>4</sub>,N<sub>24</sub>,N<sub>24</sub>-hexamethyl-, trimethiodide 125496-38-0, 2,3-Secositostane-2,3-diamine, N<sub>2</sub>,N<sub>2</sub>,N<sub>3</sub>,N<sub>3</sub>-tetramethyl-, dimethiodide (preparation of)

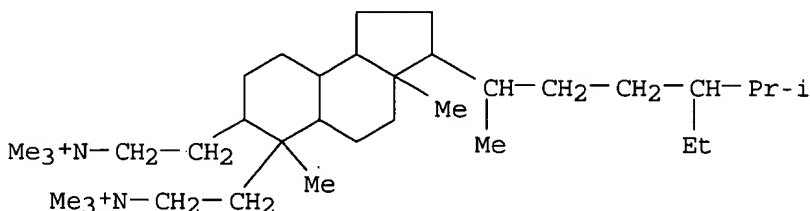
RN 122387-46-6 CAPLUS

CN 3,4-Secocholane-3,4,24-triamine, N<sub>3</sub>,N<sub>3</sub>,N<sub>4</sub>,N<sub>4</sub>,N<sub>24</sub>,N<sub>24</sub>-hexamethyl-, trimethiodide (6CI) (CA INDEX NAME)



● 3 I<sup>-</sup>

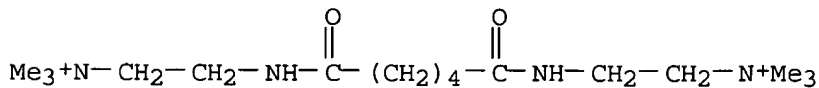
RN 125496-38-0 CAPLUS  
 CN 2,3-Secositostane-2,3-diamine, N<sub>2</sub>,N<sub>2</sub>,N<sub>3</sub>,N<sub>3</sub>-tetramethyl-, dimethiodide  
 (6CI) (CA INDEX NAME)



● 2 I<sup>-</sup>

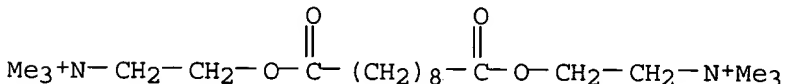
L9 ANSWER 16 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1953:70520 CAPLUS  
 DOCUMENT NUMBER: 47:70520  
 ORIGINAL REFERENCE NO.: 47:11905a-c  
 TITLE: Bolaform electrolytes. III. Conductance of bisquaternary salts of **dicarboxylic acid** bis-2-tertiary-aminoalkyl amides in methanol  
 AUTHOR(S): Eisenberg, H.; Fuoss, Raymond M.  
 CORPORATE SOURCE: Yale Univ.  
 SOURCE: Journal of the American Chemical Society (1953 ), 75, 2914-17  
 CODEN: JACSAT; ISSN: 0002-7863  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Unavailable  
 AB cf. C.A. 45, 6017e. The condens. in MeOH at 25° of the following salts were measured: N,N'-bis(2-dimethylaminoethyl)oxalamide-di-MeI, N,N'-bis(2-dimethylaminoethyl)succinamide-di-MeI, N,N'-bis(2-dimethylaminoethyl)adipamide-di-MeI, and N,N'-bis(2-dimethylaminoethyl)suberamide-di-MeI. These salts are bolaform electrolytes with, resp., 8, 10, 12, and 14 atoms joining their 2 **quaternary** nitrogens. The constant  $k_2$ , which describes the interaction of a bolaform cation and an anion, is practically the same for the oxalic and the suberic derivs., and is somewhat larger for these compds. than for the succinic and adipic derivs. This observation suggests that an intramol. ring structure, stabilized by H bonds between the **amide** groups is formed, which shortens the effective charge-charge distance.  
 IT 62055-16-7, Ammonium, [adipoylbis(iminoethylene)]bis[trimethyl-

iodide]  
 (elec. conductivity in MeOH, and structure of)  
 RN 62055-16-7 CAPLUS  
 CN Ethanaminium, 2,2'-[ (1,6-dioxo-1,6-hexanediyi)diimino]bis [N,N,N-trimethyl-, diiodide (9CI) (CA INDEX NAME)



●2 I-

L9 ANSWER 17 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1952:3443 CAPLUS  
 DOCUMENT NUMBER: 46:3443  
 ORIGINAL REFERENCE NO.: 46:626d-e  
 TITLE: The pharmacology of  $\alpha, \omega$ -bisquaternary ammonium compounds. III. Comparison of several **dicarboxylic acid esters**  
 AUTHOR(S): Ginzel, K. H.; Klupp, H.; Werner, G.  
 CORPORATE SOURCE: Univ. Vienna  
 SOURCE: Archives Internationales de Pharmacodynamie et de Therapie (1951), 87, 79-98  
 CODEN: AIPTAK; ISSN: 0003-9780  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Unavailable  
 AB The neuromuscular blocking action in nonanesthetized dogs and chloralosed cats and the spastic paralysis in pigeons decrease with increasing C-chain length of the following: bischoline esters of succinic, adipic, and sebatic acids and their ethyl derivs. The contracture response of the isolated rectus abdominis of the frog increased with chain length and was antagonized by d-tubocurarine. The hypertensive effect also increased in this manner. However, the adipic acid diester of triethyl(2-hydroxyethyl)ammonium iodide caused flaccid paralysis in pigeons and reduced the sensitivity of the frog rectus abdominis to acetylcholine.  
 IT 17140-07-7, Sebatic acid, **ester** with choline iodide (pharmacology of)  
 RN 17140-07-7 CAPLUS  
 CN Ethanaminium, 2,2'-[ (1,10-dioxo-1,10-decanediyl)bis(oxy)]bis [N,N,N-trimethyl-, diiodide (9CI) (CA INDEX NAME)



●2 I-

his

(FILE 'HOME' ENTERED AT 11:18:52 ON 27 JUN 2005)

FILE 'CAPLUS' ENTERED AT 11:19:07 ON 27 JUN 2005

L1                   STRUCTURE UPLOADED  
                  S L1

FILE 'REGISTRY' ENTERED AT 11:19:31 ON 27 JUN 2005

L2                   50 S L1

FILE 'CAPLUS' ENTERED AT 11:19:32 ON 27 JUN 2005

L3                   3 S L2  
                  S L1

FILE 'REGISTRY' ENTERED AT 11:20:12 ON 27 JUN 2005

L4                   21641 S L1 FULL

FILE 'CAPLUS' ENTERED AT 11:20:13 ON 27 JUN 2005

L5                   13121 S L4 FULL  
L6                   10617 S L5 AND PY<1999  
L7                   927 S L6 AND (ESTER OR AMIDE)  
L8                   316 S L7 AND QUATERN?  
L9                   17 S L8 AND DICARBOXYLIC ACID  
L10                  STRUCTURE UPLOADED  
                  S L1

FILE 'REGISTRY' ENTERED AT 11:29:24 ON 27 JUN 2005

L11                  50 S L1

FILE 'CAPLUS' ENTERED AT 11:29:24 ON 27 JUN 2005

L12                  3 S L11  
                  S L10

FILE 'REGISTRY' ENTERED AT 11:30:15 ON 27 JUN 2005

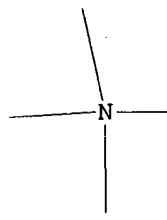
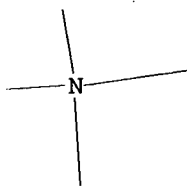
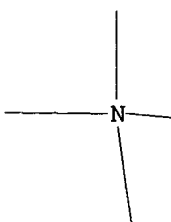
L13                  2365 S L10 FULL

FILE 'CAPLUS' ENTERED AT 11:30:16 ON 27 JUN 2005

L14                  1441 S L13 FULL  
L15                  1111 S L14 AND PY<1999  
L16                  120 S L15 AND (ESTER OR AMIDE)  
L17                  43 S L16 AND QUATERN?  
L18                  1 S L17 AND DICARBOXYLIC ACID

=>

=> d  
L10 HAS NO ANSWERS  
L10 STR



Structure attributes must be viewed using STN Express query preparation.

=> s 11  
**REG1stry INITIATED**  
Substance data SEARCH and crossover from CAS REGISTRY in progress...  
Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

SAMPLE SEARCH INITIATED 11:29:24 FILE 'REGISTRY'  
SAMPLE SCREEN SEARCH COMPLETED - 1309 TO ITERATE

76.4% PROCESSED 1000 ITERATIONS 50 ANSWERS  
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)  
SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*  
BATCH \*\*COMPLETE\*\*  
PROJECTED ITERATIONS: 24010 TO 28350  
PROJECTED ANSWERS: 21103 TO 25183

L11 50 SEA SSS SAM L1

L12 3 L11

=> d 1-3 ibib abs hitstr

L12 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN  
ACCESSION NUMBER: 2005:182792 CAPLUS  
DOCUMENT NUMBER: 142:263559  
TITLE: Preparation of shamrock surfactants and their methods  
of use  
INVENTOR(S): Jaeger, David A.  
PATENT ASSIGNEE(S): University of Wyoming, USA  
SOURCE: PCT Int. Appl., 46 pp.  
CODEN: PIXXD2  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
-----	-----	-----	-----	-----

WO 2005019405      A1      20050303      WO 2003-US29742      20030922  
 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,  
 CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE,  
 GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,  
 LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ,  
 OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM,  
 TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW  
 RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,  
 KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,  
 FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,  
 BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

PRIORITY APPLN. INFO.:      US 2003-495214P      P 20030813

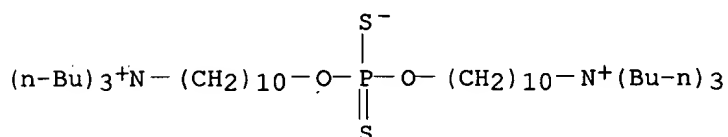
AB Shamrock surfactants are surfactants containing two ionic or polar nonionic head groups, each connected to a central ionic head group by a hydrocarbon linking moiety, wherein the central head group is a dithiophosphate, dithiocarbamate, or quaternary ammonium group. These surfactants have potential applications in chemical decontamination of mustard (simulants), storage and release devices/chemical switches and the remediation of heavy-metal ion-contaminated water. Such a surfactant compound is of the formula [X-L-Z-L'-X'] (A)p, wherein X and X' represent outer head groups, which may be the same or different and comprise charged moieties selected from the group of -N+R1R2R3, R1, R2 and R3 being the same or different, representing hydrocarbyl groups, -CO2- or -O(CH2)mSO3-, m being an integer from 2 to 30, or polar moieties of the formula, -O-(CH2CH2O)nR4, R4 being hydrogen or a C1-C6 hydrocarbyl group and n is an integer from 1 to 1000; L and L' are the same or different and represent a hydrocarbon linking moiety which may optionally be interrupted with oxygen; Z represents a central head group selected from a dithiophosphate moiety, dithiocarbamate or a quaternary ammonium moiety, wherein R5 and R6 are the same or different and represent C1-C6 hydrocarbyl groups, with the proviso that when Z represents said dithiocarbamate moiety or said quaternary ammonium moiety, X and X' do not represent N+R1R2R3, and with the further proviso that X and X' do not represent -O(CH2)mSO3- unless Z represents said quaternary ammonium moiety; and A represents a counter ion, which may be either pos. or neg. depending on the net charge of [X-L-Z-L'-X'] and p is an integer which, when multiplied by the valency of said counter ion yields the absolute value of the net charge of [X-L-Z-L'-X'].

IT 813446-48-9P

RL: PRP (Properties); SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses)  
 (preparation of shamrock surfactants and their methods of use)

RN 813446-48-9 CAPLUS

CN 16,18-Dioxa-5-azonia-17-phosphaoctacosan-28-aminium, N,N,N,5,5-pentabutyl-17-mercaptop-, inner salt, bromide, 17-sulfide (9CI) (CA INDEX NAME)



● Br<sup>-</sup>

REFERENCE COUNT:      1      THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

DOCUMENT NUMBER: 142:446807  
 TITLE: Promotion effect of cationic Gemini surfactants on  
 1-dodecene hydroformylation in biphasic catalytic  
 system  
 AUTHOR(S): Xu, Bin; Li, Min; Yang, Min; Zheng, Hong-Jie; He,  
 Yu-E.; Chen, Hua; Li, Xian-Jun  
 CORPORATE SOURCE: Institute of Homogeneous catalysis, College of  
 Chemistry, Sichuan University, Chengdu, 610064, Peop.  
 Rep. China  
 SOURCE: Gaodeng Xuexiao Huaxue Xuebao (2004), 25(11),  
 2060-2064  
 CODEN: KTHPDM; ISSN: 0251-0790  
 PUBLISHER: Gaodeng Jiaoyu Chubanshe  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Chinese

**AB** New cationic Gemini surfactants with a rigid spacer group of xylene were synthesized, their compns. and structures were characterized. The cmc and the solubilizations of 1-dodecene in micellar solution were determined by surface

tension method and UV-Vis spectrometry resp. The cmc of new Gemini surfactants are lower than CTAB by about an order, but the solubilizations of 1-dodecene in Gemini surfactants solution are higher than that in CTAB solution. The promotion effect of cationic Gemini surfactants on 1-dodecene hydroformylation in biphasic catalytic system was studied. The results indicated that in the biphasic system containing the catalysts RhCl(CO)(TPPTS)2 and TPPTS, [TPPTS = tris (sodium-m-sulfonatophenyl) phosphine], the reaction rate of 1-dodecene hydroformylation in the presence of Gemini surfactants was faster than that in the presence of conventional surfactant CATB, e.g., the conversion of 1-dodecene is 90% when G (o-xyl)c22 concentration in aqueous solution was 2 + 10-3 mol/L,

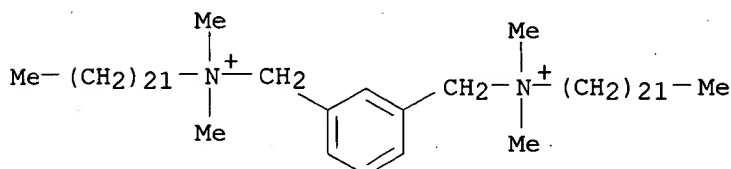
however, the conversion was only 20.4% when the concentration of CTAB was 3 + 10-3 mol/L. The greater acceleration of Gemini surfactants in 1-dodecene hydroformylation could be attributed to that cationic Gemini surfactants had lower cmc and better solubilization for the substrate. The lower cmc and surface tension are favorable for increasing the interfacial area of two phases, breaking phase barrier and promoting the substrate transfer to interface where the substrates coordinate with the active rhodium complex anion species. The regioselectivity for olefin hydroformylation in Gemini surfactants.

IT 851319-26-1

RL: NUU (Other use, unclassified); PRP (Properties); RGT (Reagent); RACT (Reactant or reagent); USES (Uses)  
 (promotion effect of cationic Gemini surfactants on 1-dodecene hydroformylation in biphasic catalytic system)

RN 851319-26-1 CAPLUS

CN 1,3-Benzenedimethanaminium, N,N'-didocosyl-N,N,N',N'-tetramethyl-, dibromide (9CI) (CA INDEX NAME)

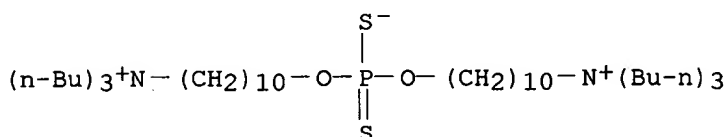


L12 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 2004:902647 CAPLUS  
 DOCUMENT NUMBER: 142:76557  
 TITLE: Shamrock Surfactants: Synthesis and Characterization  
 AUTHOR(S): Jaeger, David A.; Zeng, Xiaohui; Apkarian, Robert P.  
 CORPORATE SOURCE: Department of Chemistry, University of Wyoming,  
 Laramie, WY, 82071, USA  
 SOURCE: Langmuir (2004), 20(24), 10427-10432  
 CODEN: LANGD5; ISSN: 0743-7463  
 PUBLISHER: American Chemical Society  
 DOCUMENT TYPE: Journal  
 LANGUAGE: English

**AB** Two types of a new class of surfactants with 3 headgroups were prepared. A central headgroup is connected to 2 flanking headgroups by hydrocarbon chains. The term "shamrock" is used to describe these surfactants, denoting their triple-headed character and reflecting the fact that shamrocks have leaflets in groups of 3. The major lipophilic character of shamrock surfactants is provided by the 2 hydrocarbon chains linking the 3 headgroups and not by long-chain alkyl groups appended to the linking hydrocarbon chains or the headgroups. The new surfactants are (2,2,15,15,28,28-hexamethyl-2,15,28-triaazonianonacosane triiodide), (2,2,15,15,28,28-hexamethyl-2,15,28-triaazonianonacosane trichloride) (I), (O,O'-di-[10-(N,N,N-tripropylammonio)decyl]phosphorodithioate bromide), and (O,O'-di-[10-(N,N,N-tributylammonio)decyl]phosphorodithioate bromide). (2,2,9,9,16,16-Hexamethyl-2,9,16-triaazoniaheptadecane triiodide) was prepared for comparison. The surfactants were characterized in water by measurement of their Krafft temps. and critical aggregation concns., and their aggregates were studied by  $^1\text{H}$  NMR spectroscopy, dynamic laser light scattering, and phase-contrast optical microscopy. Aqueous I was also studied by cryo-etch high-resolution SEM, which revealed irregularly shaped cells containing a complex matrix of surfactant. Coacervates were observed by optical microscopy upon the hydration of the shamrock surfactants.

**IT** 813446-48-9P  
 RL: PRP (Properties); SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses)  
 (synthesis and characterization of trilobal shamrock surfactants)

**RN** 813446-48-9 CAPLUS  
**CN** 16,18-Dioxa-5-azonia-17-phosphaoctacosan-28-aminium, N,N,N,5,5-pentabutyl-17-mercaptop-, inner salt, bromide (9CI) (CA INDEX NAME)



● Br<sup>-</sup>

REFERENCE COUNT: 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS  
 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s 110 full

REGISTRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress...  
 Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SEARCH INITIATED 11:30:15 FILE 'REGISTRY'  
FULL SCREEN SEARCH COMPLETED - 3731 TO ITERATE

100.0% PROCESSED 3731 ITERATIONS ( 2 INCOMPLETE) 2365 ANSWERS  
SEARCH TIME: 00.00.01

L13 2365 SEA SSS FUL L10

L14 1441 L13

=> s 114 and py<1999  
18930685 PY<1999

L15 1111 L14 AND PY<1999

=> s 115 and (ester or amide)  
562887 ESTER  
119577 AMIDE  
L16 120 L15 AND (ESTER OR AMIDE)

=> s 116 and quatern?  
133137 QUATERN?  
L17 43 L16 AND QUATERN?

=> s 117 and dicarboxylic acid  
60801 DICARBOXYLIC  
3995178 ACID  
36503 DICARBOXYLIC ACID  
(DICARBOXYLIC(W)ACID)  
L18 1 L17 AND DICARBOXYLIC ACID

=> d ibib abs hitstr

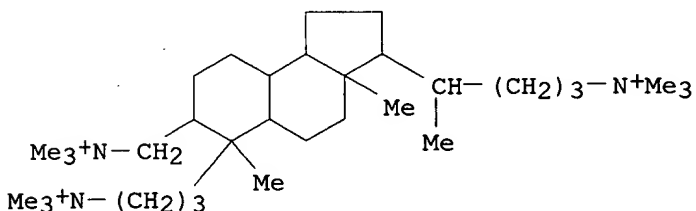
L18 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2005 ACS on STN  
ACCESSION NUMBER: 1958:50718 CAPLUS  
DOCUMENT NUMBER: 52:50718  
ORIGINAL REFERENCE NO.: 52:9173i,9174a-c  
TITLE: Multivalent **quaternary** ammonium compounds.  
VI. Some reaction products of bile acids and sterols  
AUTHOR(S): Lettre, H.; Gottstein, W.; Scholtissek, Ch.  
CORPORATE SOURCE: Univ. Heidelberg, Germany  
SOURCE: Monatshefte fuer Chemie (1957), 88, 715-20  
CODEN: MOCMB7; ISSN: 0026-9247  
DOCUMENT TYPE: Journal  
LANGUAGE: Unavailable  
AB cf. C.A. 51, 4409a. Some N derivs. of lithiobilanic acid (I) and  
sitosterol are prepared I is treated with Ac<sub>2</sub>O followed by Me<sub>2</sub>NH to yield  
lithiobilanic acid 3-monodimethylamide, m. 251-2°. I with Ac<sub>2</sub>O  
followed by PC<sub>1</sub>5 and then with Me<sub>2</sub>NH in Et<sub>2</sub>O gives an Et<sub>2</sub>O phase containing  
60%-70% I 3,4,24-tris(dimethylamide), m. 151-2°, purified by  
chromatography on Al<sub>2</sub>O<sub>3</sub>. The aqueous phase of the reaction yields 15-20% of I  
3,24-bis(dimethylamide) (II), m. 232-3°. II is esterified with  
CH<sub>2</sub>N<sub>2</sub> and reduced with LiAlH<sub>4</sub> in tetrahydrofuran to 3,4-secocholan-4-ol-  
3,24-bis(dimethylamine hydrochloride), m. 292-5° (decomposition). II is  
similarly reduced to 90% 3,4-secocholane-3,4,24-tris(dimethylamine  
hydrochloride), decompose 275°, which forms 3,4-secocholane-3,24-

tris(trimethylammonium iodide), m. 290° (decomposition). The **dicarboxylic acid** of sitosterol (III), heated 2 hrs. with Ac<sub>2</sub>O gives 76% 2,3-secositostanol-2,3-**dicarboxylic acid** anhydride, m. 176°. III di-Me **ester** is reduced by LiAlH<sub>4</sub> to 88% 2,3-secositostane-2,3-diol, m. 182-3° (MeOH). III with PCl<sub>5</sub> and Me<sub>2</sub>NH yields by chromatography on Al<sub>2</sub>O<sub>3</sub> 48% 2,3-secositostane-2,3-**dicarboxylic acid** dimethylamide, m. 106-7°, reduced by LiAlH<sub>4</sub> to 68% 2,3-secositostane-2,3-bis(dimethylamine hydrochloride), m. 326° (decomposition). This compound with MeI gives 2,3-secositostane-2,3-bis(trimethylammonium iodide), m. 323°.

IT 122387-46-6, 3,4-Secocholane-3,4,24-triamine, N<sub>3</sub>,N<sub>3</sub>,N<sub>4</sub>,N<sub>4</sub>,N<sub>24</sub>,N<sub>24</sub>-hexamethyl-, trimethiodide  
(preparation of)

RN 122387-46-6 CAPLUS

CN 3,4-Secocholane-3,4,24-triamine, N<sub>3</sub>,N<sub>3</sub>,N<sub>4</sub>,N<sub>4</sub>,N<sub>24</sub>,N<sub>24</sub>-hexamethyl-, trimethiodide (6CI) (CA INDEX NAME)



●3 I-

=> d 117 1-10 ibib abs hitstr

L17 ANSWER 1 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2002:737708 CAPLUS

DOCUMENT NUMBER: 137:237406

TITLE: A **quaternary** ammonium phosphate-containing aqueous composition suitable for the application to human skin

INVENTOR(S): Zeigler, Philip Dale; Cheney, Michael Charles

PATENT ASSIGNEE(S): Hindustan Lever Ltd., India

SOURCE: Indian, 40 pp.

CODEN: INXXAP

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
IN 173884	A	19940730	IN 1992-B05691	19920220 <--
PRIORITY APPLN. INFO.:			IN 1992-B05691	19920220

OTHER SOURCE(S): MARPAT 137:237406

AB An aqueous composition is provided which includes a **quaternary** ammonium functionalized phosphate **ester** and a cationic polysaccharide. The compns. may include an emollient, and be used as a water-proof sunscreen composition. For example, a sunscreen composition was prepared from

(by

weight): Phase A containing cetyl alc. 2.5%, glycerol monostearate (Kessco GMS) 1.5%, Pr paraben 0.10%, ethylhexyl p-methoxycinnamate (Parsol MCX) 7.0%, oxybenzone (Uvinul M-40) 3.0%, octyl palmitate (Schercemol OP) 2.0%, silicone fluid 1.0%, and petroleum jelly 1.0%, and Phase B containing glycerin 4.0%, Monaquat P-TS 3.0%, Antifoam AF 0.005%, Me paraben 0.150%, Quatrisoft LM-200 0.250%, fragrance 0.150%, and water up to 100%.

IT 144377-73-1, Phospholipid EFA 144379-29-3, Monaquat P-TS

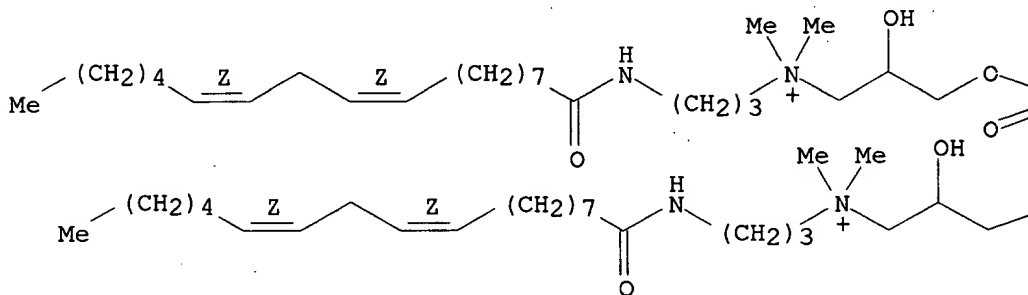
RL: COS (Cosmetic use); BIOL (Biological study); USES (Uses)  
(cosmetic compns. containing **quaternary** ammonium phosphate and cationic polysaccharide)

RN 144377-73-1 CAPLUS

CN 4,6-Dioxa-14-aza-10-azonia-5-phosphadotriaconta-23,26-dien-1-aminium, 5-[3-[dimethyl[3-[(9Z,12Z)-1-oxo-9,12-octadecadienyl]amino]propyl]ammonio]-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-15-oxo-N-[3-[(9Z,12Z)-1-oxo-9,12-octadecadienyl]amino]propyl]-, trichloride, 5-oxide, (23Z,26Z)- (9CI) (CA INDEX NAME)

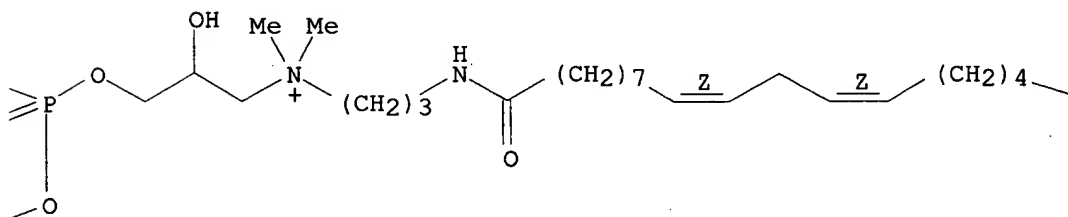
Double bond geometry as shown.

PAGE 1-A



● 3 Cl<sup>-</sup>

PAGE 1-B



PAGE 1-C

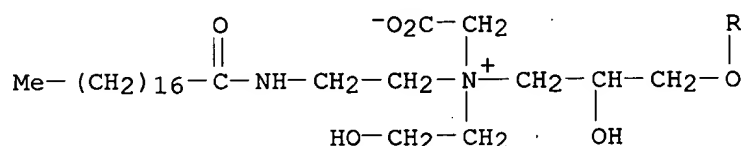
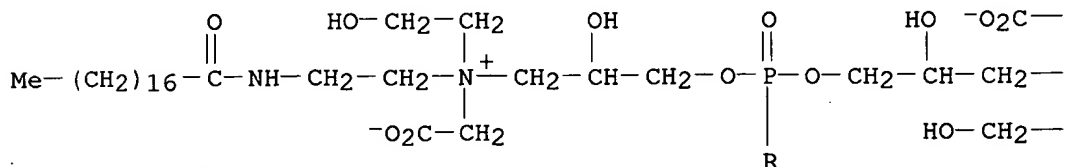
Me

RN 144379-29-3 CAPLUS

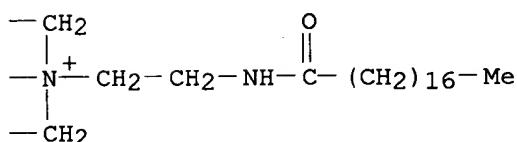
CN 4,6-Dioxa-13-aza-10-azonia-5-phosphahentriacontan-1-aminium,

N,10-bis (carboxymethyl)-5-[3-[(carboxymethyl)(2-hydroxyethyl)[2-[(1-oxooctadecyl)amino]ethyl]ammonio]-2-hydroxypropoxy]-2,8-dihydroxy-N,10-bis(2-hydroxyethyl)-14-oxo-N-[2-[(1-oxooctadecyl)amino]ethyl]-, tris(inner salt), 5-oxide (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 1-B



L17 ANSWER 2 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
ACCESSION NUMBER: 1998:766507 CAPLUS  
DOCUMENT NUMBER: 130:29221  
TITLE: Preparation of solid porous matrixes for pharmaceutical uses  
INVENTOR(S): Unger, Evan C.  
PATENT ASSIGNEE(S): ImaRx Pharmaceutical Corp., USA  
SOURCE: PCT Int. Appl., 139 pp.  
CODEN: PIXXD2  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 6  
PATENT INFORMATION:

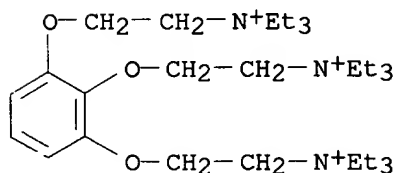
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9851282	A1	19981119	WO 1998-US9570	19980512 <--
W: AU, BR, CA, CN, JP, KR, NZ				
RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,				
PT, SE				
US 2002039594	A1	20020404	US 1998-75477	19980511
AU 9873787	A1	19981208	AU 1998-73787	19980512 <--
EP 983060	A1	20000308	EP 1998-921109	19980512
R: DE, FR, GB, IT, NL				
US 2001018072	A1	20010830	US 2001-828762	20010409
US 2004091541	A1	20040513	US 2003-622027	20030716
PRIORITY APPLN. INFO.:			US 1997-46379P	P 19970513
			US 1998-75477	A 19980511

AB A solid porous matrix formed from a surfactant, a solvent, and a bioactive agent is described. Thus, amphotericin nanoparticles were prepared by using ZrO<sub>2</sub> beads and a surfactant. The mixture was milled for 24 h.

IT 65-29-2, Gallamine triethiodide  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (preparation of solid porous matrixes for pharmaceutical uses)

RN 65-29-2 CAPLUS

CN Ethanaminium, 2,2',2'''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-, triiodide (9CI) (CA INDEX NAME)



●3 I-

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 3 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1997:648717 CAPLUS  
 DOCUMENT NUMBER: 127:333127  
 TITLE: Softening agents containing polycationic surfactants for fabrics in laundering  
 INVENTOR(S): Imada, Hiroshi; Imai, Hiroto; Sugafuji, Hisahiro;  
 Fujiwara, Masami  
 PATENT ASSIGNEE(S): Lion Corp., Japan  
 SOURCE: Jpn. Kokai Tokkyo Koho, 10 pp.  
 CODEN: JKXXAF  
 DOCUMENT TYPE: Patent  
 LANGUAGE: Japanese  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 09256273	A2	19970930	JP 1996-67691	19960325 <--
PRIORITY APPLN. INFO.:			JP 1996-67691	19960325

OTHER SOURCE(S): MARPAT 127:333127

AB The agents, used during the rinse cycle, which soften fabrics without affecting hydrophilicity and yellowing prevention, contain polycationic surfactants R<sub>3</sub>(R<sub>1</sub>R<sub>4</sub>N<sup>+</sup>A)<sub>n</sub>N<sup>+</sup>R<sub>2</sub>R<sub>5</sub>R<sub>6</sub> (n + 1)X<sup>-</sup>, cationic surfactants R<sub>7</sub>(N<sup>+</sup>R<sub>8</sub>R<sub>9</sub>G)<sub>m</sub>N<sup>+</sup>R<sub>10</sub>R<sub>11</sub>R<sub>12</sub> (m + 1)X<sup>-</sup>, and anionic surfactants R<sub>13</sub>CH(SO<sub>3</sub>X<sub>1</sub>)CO<sub>2</sub>R<sub>14</sub> (A = C<sub>2</sub>-12 alkylene, hydroxyalkylene; G = C<sub>2</sub>-10 alkylene; R<sub>1</sub>, R<sub>2</sub>, R<sub>7</sub> = C<sub>10</sub>-28 saturated hydrophobic group; R<sub>3</sub>-R<sub>6</sub>, R<sub>8</sub>-R<sub>12</sub> = C<sub>1</sub>-6 alkyl, hydroxyalkyl; R<sub>13</sub> = C<sub>8</sub>-26 alkyl; R<sub>14</sub> = C<sub>1</sub>-6 alkyl; X = halogen; X<sub>1</sub> = H, metal, ammonium; m ≥ 0; n ≥ 1). Thus, reacting 2 mol N,N-dimethylstearylamine and 1 mol 1,6-diiodohexane to give a diquaternary ammonium salt (I), sep. reacting 1 mol Duomeen HT Flake, 4 mol HCHO, and excess HCO<sub>2</sub>H, followed by reaction with MeCl to give another quaternary ammonium salt (II), mixing I 0.9, II 0.1, and Na Me α-sulfostearate 0.5 equiv, and dissolving the mixture in water at 3%

gave a softening agent. A cotton towel was washed and rinsed in a washing machine; then the agent was added at 0.0033%. After the towel was squeezed and dried, it showed good softness and good water absorbency.

IT 197862-15-0P

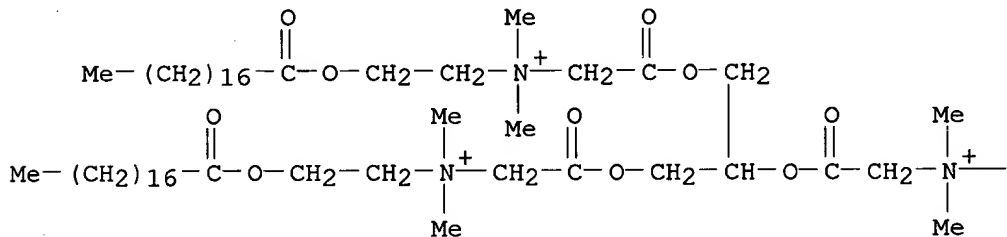
RL: IMF (Industrial manufacture); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses)

(polycationic; softening agents containing polycationic surfactants, cationic surfactants, and anionic surfactants for fabrics after laundering)

RN 197862-15-0 CAPLUS

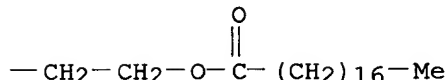
CN 3,7,13-Trioxa-10-azoniahetriacantan-1-aminium, 5-[[[dimethyl[2-[(1-oxooctadecyl)oxy]ethyl]ammonio]acetyl]oxy]-N,N,10,10-tetramethyl-2,8,14-trioxa-N-[2-[(1-oxooctadecyl)oxy]ethyl]-, trichloride (9CI) (CA INDEX NAME)

PAGE 1-A



● 3 Cl-

PAGE 1-B



L17 ANSWER 4 OF 43 CAPIUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1997:369695 CAPLUS

DOCUMENT NUMBER: 126:347155

TITLE: Cosmetic compositions containing cationic resin and waxes

INVENTOR(S): Sheard, Christine

PATENT ASSIGNEE(S): Boots Company Plc, UK; Sheard, Christine

SOURCE: PCT Int. Appl., 21 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

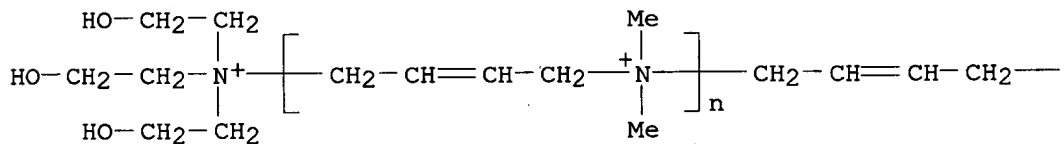
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9713497	A1	19970417	WO 1996-EP4393	19961009 <-- W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE,

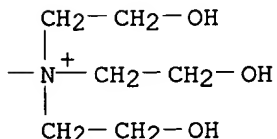
ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS,  
 LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD,  
 SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, AM, AZ, BY,  
 KG, KZ, MD, RU, TJ, TM  
 RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR,  
 IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA  
 AU 9672893 A1 19970430 AU 1996-72893 19961009 <--  
 EP 862410 A1 19980909 EP 1996-934606 19961009 <--  
 EP 862410 B1 20021218  
 R: DE, FR, GB  
 ZA 9608552 A 19980610 ZA 1996-8552 19961010 <--  
 GB 1995-20690 A 19951010  
 WO 1996-EP4393 W 19961009  
 PRIORITY APPLN. INFO.:  
 AB A cosmetic composition comprises 0.05-5% hydrophilic cationic resin 30-85% oil component 1-40% wax component and 1-40 % weight/weight powder component. The hydrophilic cationic resin may be water soluble or water swellable and may also be any mixture of suitable homopolymers or copolymers, e.g., any mixture of 1 or more Polyquaternium polymers or polymeric salts preferably those denoted by the CFTA name Polyquaternium. The cosmetic composition is solid at ambient temperature and is suitable for use as a lipstick. A product comprising  
 the composition associated with a suitable receptacle and/or dispenser is also disclosed. Thus, a lipstick contained plant wax 6.4, paraffin wax 9.0, synthetic wax 2.3, synthetic fat 10.0, fatty alc. 20.7, synthetic ester 12.74, plant oil 26.24, preservative 0.1, antioxidant 0.03, Salcare SC96 2.25, butylene glycol 1.5, and pigment 8.74%.  
 IT 75345-27-6, Polyquaternium-1  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (cosmetic compns. containing cationic resin and waxes)  
 RN 75345-27-6 CAPLUS  
 CN Poly[(dimethyliminio)-2-butene-1,4-diyl chloride],  $\alpha$ -[4-[tris(2-hydroxyethyl)ammonio]-2-butenyl]- $\omega$ -[tris(2-hydroxyethyl)ammonio]-, dichloride (9CI) (CA INDEX NAME)

PAGE 1-A



●3 Cl<sup>-</sup>

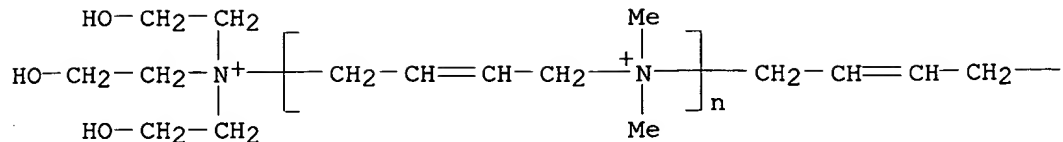
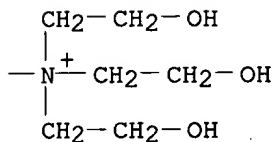
PAGE 1-B



TITLE: Body wash compositions containing anionic cleansing surfactants polymeric cationic conditioning compounds and **quaternized** phosphate esters  
 INVENTOR(S): Scafidi, Anthony A.  
 PATENT ASSIGNEE(S): Helene Curtis, Inc., USA  
 SOURCE: PCT Int. Appl., 59 pp.  
 CODEN: PIXXD2  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9710804	A1	19970327	WO 1996-US14410	19960909 <--
W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, UZ, VN, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI				
US. 5683683	A	19971104	US 1995-531712	19950921 <--
ZA 9607294	A	19970304	ZA 1996-7294	19960828 <--
CA 2231809	AA	19970327	CA 1996-2231809	19960909 <--
AU 9669697	A1	19970409	AU 1996-69697	19960909 <--
BR 9610522	A	19990706	BR 1996-10522	19960909
PRIORITY APPLN. INFO.:			US 1995-531712	A 19950921
			WO 1996-US14410	W 19960909

OTHER SOURCE(S): MARPAT 126:308638  
 AB A body wash composition containing an anionic cleansing surfactant, such as an alkyl ether sulfate or an alkyl sulfate, like sodium lauryl ether sulfate or sodium lauryl sulfate; a polymeric cationic conditioning compound, such as a **quaternized** guar gum; and a **quaternized** phosphate ester in an aqueous carrier is disclosed. The composition is used to cleanse and to impart conditioning properties to the skin. A body wash composition contained sodium lauryl ether sulfate 12.0, a premixed surfactant concentrate 3.6, cocamide MEA 7.0, preservatives 0.5, guar hydroxypropyltrimonium chloride 0.2, tetrasodium ethylenediamine tetraacetic acid 0.08, citric acid 0.15, palmitic acid 2.0, stearamidopropyl phosphatidyl PG-dimonium chloride 0.4, cocamidopropyl hydroxysultate 1.9, titanium dioxide 0.2, and water q.s. 100%.  
 IT 75345-27-6, Polyquaternium 1  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (body wash compns. containing anionic cleansing surfactants polymeric cationic conditioning compds. and **quaternized** phosphate esters)  
 RN 75345-27-6 CAPLUS  
 CN Poly[(dimethyliminio)-2-butene-1,4-diyl chloride],  $\alpha$ -[4-[tris(2-hydroxyethyl)ammonio]-2-butenyl]- $\omega$ -[tris(2-hydroxyethyl)ammonio]-, dichloride (9CI) (CA INDEX NAME)

● 3 Cl<sup>-</sup>

L17 ANSWER 6 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1996:656984 CAPLUS  
 DOCUMENT NUMBER: 125:308699  
 TITLE: Emulsified, low pH cosmetic compositions having improved stability  
 INVENTOR(S): Papadakis, Marcelline C.  
 PATENT ASSIGNEE(S): Helene Curtis, Inc., USA  
 SOURCE: U.S., 10 pp.  
 CODEN: USXXAM  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5567427	A	19961022	US 1995-406106	19950317 <--
WO 9629051	A1	19960926	WO 1996-US3761	19960314 <--
W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI				
RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML				
AU 9653175	A1	19961008	AU 1996-53175	19960314 <--
PRIORITY APPLN. INFO.: US 1995-406106 A 19950317 WO 1996-US3761 W 19960314				

AB Emulsified, low pH cosmetic compns. having improved pH and phase stability are disclosed. The emulsified cosmetic compns. have a pH 3.7-4.5, and contain 10-50% by weight dispersed oil phase, 2-20% by weight acid, like a hydroxycarboxylic acid, and 0.5-2% **quaternized** phosphate ester, like linoleamidopropyl PG-dimonium chloride phosphate. The emulsified cosmetic compns. are phase-stable over an extended storage period and maintain a constant pH by exhibiting a pH drift of about 0.15, usually  $\leq 0.1$  pH units.

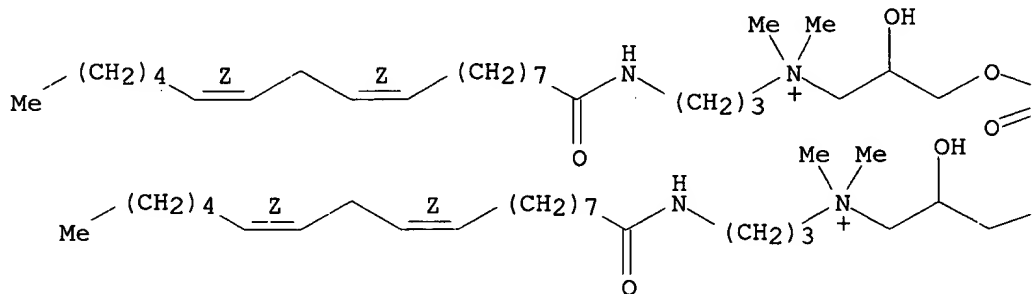
IT 144377-73-1, Phospholipid EFA  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (emulsified, stable, low pH cosmetic compns. containing)

RN 144377-73-1 CAPLUS

CN 4,6-Dioxa-14-aza-10-azonia-5-phosphadotriaconta-23,26-dien-1-aminium,  
 5-[3-[dimethyl[3-[(9Z,12Z)-1-oxo-9,12-octadecadienyl]amino]propyl]ammonio  
 ]-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-15-oxo-N-[3-  
 [(9Z,12Z)-1-oxo-9,12-octadecadienyl]amino]propyl]-, trichloride, 5-oxide,  
 (23Z,26Z)- (9CI) (CA INDEX NAME)

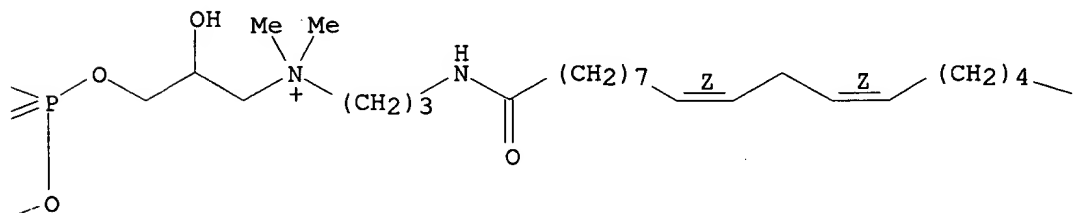
Double bond geometry as shown.

PAGE 1-A



●3 Cl<sup>-</sup>

PAGE 1-B



PAGE 1-C

Me

L17 ANSWER 7 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1996:377496 CAPLUS  
 DOCUMENT NUMBER: 125:118113  
 TITLE: Automatic dishwashing detergent compositions comprising multiquaternary bleach activators  
 INVENTOR(S): Sivik, Mark R.; Taylor, Lucille F.; Burckett St.Laurent, James C. T.  
 PATENT ASSIGNEE(S): The Procter and Gamble Company, USA  
 SOURCE: U.S., 20 pp., Cont.-in-part of U.S. Ser. No. 298,904.  
 CODEN: USXXAM  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5520835	A	19960528	US 1995-438126	19950508 <--
US 5578136	A	19961126	US 1994-298904	19940831 <--
US 5654421	A	19970805	US 1995-486654	19950607 <--
CA 2154704	AA	19960301	CA 1995-2154704	19950726 <--
CA 2154704	C	19990615		
CA 2244021	AA	19960301	CA 1995-2244021	19950726 <--
CA 2244021	C	20021119		
ES 2202342	T3	20040401	ES 1995-305458	19950804
EP 742280	A2	19961113	EP 1996-302491	19960409 <--
EP 742280	A3	19991201		
R: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE				
CA 2175275	AA	19961109	CA 1996-2175275	19960429 <--
CA 2175275	C	19990831		
PRIORITY APPLN. INFO.:			US 1994-298904	A2 19940831
			US 1995-438126	A 19950508

OTHER SOURCE(S):

OTHER SOURCE(S): MARPAT 125:118113

AB. The title compounds, especially granular detergents, comprise bleach and

AB The title compound, especially granular detergent, contains bleaching agent structures specified) containing multiple **quaternary** N groups, preferably  $\geq 3$  such groups and preferably have  $\geq 1$  **quaternary** N group in the peracid-forming portion of the bleach activator as well as  $\geq 1$  **quaternary** N group in the leaving-group portion. Thus,  $\text{H}_2\text{N}(\text{CH}_2)_5\text{CO}_2\text{H}\cdot\text{HCl}$  obtained by hydrolysis of  $\epsilon$ -caprolactam with HCl was N-methylated with HCHO/HCO<sub>2</sub>H, the product was acid chlorinated with COCl<sub>2</sub>, then esterified with HOCH(CH<sub>2</sub>NMe<sub>2</sub>)<sub>2</sub> and the **ester quaternized** with MeCl to give the title activator 3-[Me<sub>3</sub>N<sup>+</sup>(CH<sub>2</sub>)<sub>5</sub>CO<sub>2</sub>]CH(CH<sub>2</sub>N<sup>+</sup>Me<sub>3</sub>)<sub>2</sub>·3HCl.

IT 179325-36-1DP, salts with compatible anions 179325-37-2P

179325-38-3P 179325-39-4DP, salts with compatible anions

179325-40-7DP, salts with compatible anions 179325-41-8DP

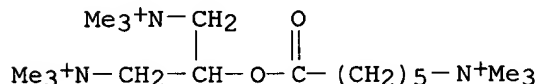
, salts with compatible anions

RL: IMF (Industrial manufacture); PREP (Preparation)

(automatic dishwashing detergent compns. comprising multiquaternary bleach activators)

RN 179325-36-1 CAPLUS

CN 1,3-Propanediaminium, N,N,N,N',N',N'-hexamethyl-2-[(1-oxo-6-(trimethylammonio)hexyl]oxy]- (9CI) (CA INDEX NAME)



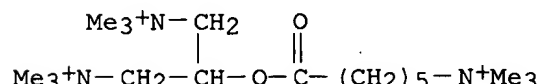
RN 179325-37-2 CAPLUS

CN 1,3-Propanediaminium, N,N,N,N',N',N'-hexamethyl-2-[(1-oxo-6-(trimethylammonio)hexyl]oxy]-, tris(methyl sulfate) (9CI) (CA INDEX NAME)

CM 1

CRN 179325-36-1

CMF C18 H42 N3 O2

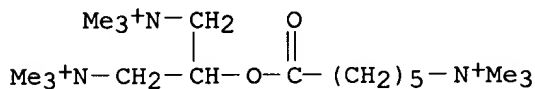


CM 2

CRN 21228-90-0  
CMF C H3 O4 S

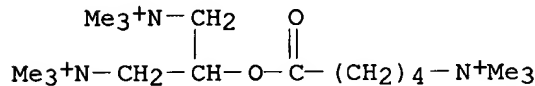
Me-O-SO<sub>3</sub><sup>-</sup>

RN 179325-38-3 CAPLUS  
CN 1,3-Propanediaminium, N,N,N,N',N',N'-hexamethyl-2-[[1-oxo-6-(trimethylammonio)hexyl]oxy]-, trichloride (9CI) (CA INDEX NAME)

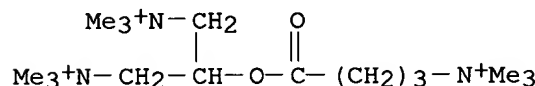


● 3 Cl<sup>-</sup>

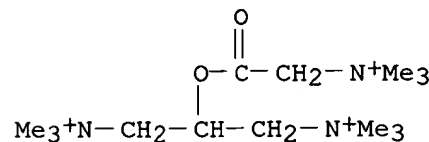
RN 179325-39-4 CAPLUS  
CN 1,3-Propanediaminium, N,N,N,N',N',N'-hexamethyl-2-[[1-oxo-5-(trimethylammonio)pentyl]oxy]- (9CI) (CA INDEX NAME)



RN 179325-40-7 CAPLUS  
CN 1,3-Propanediaminium, N,N,N,N',N',N'-hexamethyl-2-[[1-oxo-4-(trimethylammonio)butoxy]- (9CI) (CA INDEX NAME)



RN 179325-41-8 CAPLUS  
CN 1,3-Propanediaminium, N,N,N,N',N',N'-hexamethyl-2-[[[(trimethylammonio)acetyl]oxy]- (9CI) (CA INDEX NAME)



L17 ANSWER 8 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
ACCESSION NUMBER: 1996:153898 CAPLUS  
DOCUMENT NUMBER: 124:264081  
TITLE: Liquid bleaching compositions for textiles with good

INVENTOR(S): storage stability  
 Ogura, Nobuyuki; Ozaki, Kazuyoshi; Hishige, Takaomi;  
 Aoyanagi, Muneo  
 Kao Corp, Japan  
 SOURCE: Jpn. Kokai Tokkyo Koho, 13 pp.  
 CODEN: JKXXAF

DOCUMENT TYPE: Patent  
 LANGUAGE: Japanese  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 07331289	A2	19951219	JP 1994-129113	19940610 <--
JP 3330226	B2	20020930		
			JP 1994-129113	19940610

PRIORITY APPLN. INFO.: MARPAT 124:264081

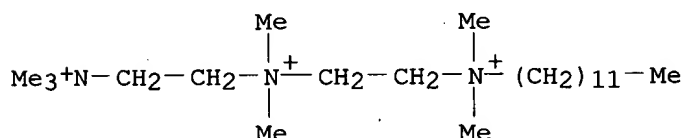
OTHER SOURCE(S): AB The bleaching compns. contain H<sub>2</sub>O<sub>2</sub>, bleaching activators which form organic acids on reaction with H<sub>2</sub>O<sub>2</sub>, polycationic compds. R<sub>1</sub>R<sub>2</sub>R<sub>3</sub>N<sup>+</sup>(R<sub>4</sub>N<sup>+</sup>R<sub>5</sub>R<sub>6</sub>)<sub>n</sub>R<sub>7</sub>. (n + 1)Z- [R<sub>1</sub>, R<sub>2</sub>, R<sub>3</sub>, R<sub>5</sub>, R<sub>6</sub>, and/or R<sub>7</sub> is C<sub>8</sub>-22 alkyl, alkenyl, (C<sub>1</sub>-22 alkyl-substituted) aryl and the remainder is (OH-containing) C<sub>1</sub>-4 alkyl; R<sub>4</sub> = ester, amido, (OH-containing) C<sub>2</sub>-6 alkylene; n = 1-3; Z-1 = anion], and nonionic surfactants, amphoteric surfactants, and/or anionic surfactants. Thus, 5% H<sub>2</sub>O<sub>2</sub>, 1% Me(CH<sub>2</sub>)<sub>10</sub>COO-1,4-C<sub>6</sub>H<sub>4</sub>SO<sub>3</sub>Na, 2% C<sub>12</sub>H<sub>25</sub>Me<sub>2</sub>N+C<sub>2</sub>H<sub>4</sub>N+Me<sub>3</sub>.2Cl-, 5% C<sub>12</sub>H<sub>25</sub>Me<sub>2</sub>N+CH<sub>2</sub>CH(OH)CH<sub>2</sub>.SO<sub>3</sub>-, 0.1% (HO)<sub>2</sub>P(O)C(OH)MeP(O)(OH)<sub>2</sub>, and balance H<sub>2</sub>O were mixed to give a composition exhibiting bleaching activity retention 97.3% after 5 mo at 50° and 80% relative humidity.

IT 175539-18-1P

RL: IMF (Industrial manufacture); MOA (Modifier or additive use); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses) (additive; for liquid bleaching compns. containing hydrogen peroxide for textiles with good storage stability)

RN 175539-18-1 CAPLUS

CN 1,2-Ethanediuminium, N-[2-(dodecyldimethylammonio)ethyl]-N,N,N',N',N'-pentamethyl-, trichloride (9CI) (CA INDEX NAME)



●3 Cl-

L17 ANSWER 9 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1994:704341 CAPLUS  
 DOCUMENT NUMBER: 121:304341  
 TITLE: Gas chromatographic separation of linear hydrocarbons on microporous organo-smectites  
 AUTHOR(S): Lao, Hongbai; Detellier, Christian  
 CORPORATE SOURCE: Ottawa-Carleton Chemistry Institute, University of Ottawa, Ottawa, ON, K1N 6N5, Can.  
 SOURCE: Clays and Clay Minerals (1994), 42(4), 477-81  
 CODEN: CLCMAB; ISSN: 0009-8604  
 PUBLISHER: Clay Minerals Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB A series of organo-montmorillonites and organo-hectorites were prepared by complete ion-exchange from the pure sodium form of the parent smectites.

The organic cations were tetramethylammonium, tri-Me **quaternary** ammonium derivs. of lysine Me **ester** and ornithine Me **ester**, **quaternized** polyammonium cations, or tetraphenylphosphonium (TPP). These organo-smectites were used as packing material for gas chromatog. columns. Mixts. of light (C1-4) hydrocarbons could be separated. The degree of separation depends on the presence of micropores

or of organophilic mesopores. The BET surface area, the micropore and mesopore vols., as well as the size distribution of micropores and mesopores, were measured for several systems. As a general trend, the retention times of the light hydrocarbons decreased with increasing micropore volume. In the case of TPP-montmorillonite, characterized by a large mesopore volume but for which no microporosity could be detected, separation of longer (C5-8) alkanes could also be achieved.

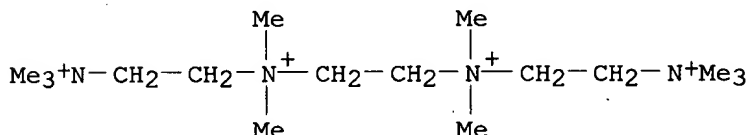
IT 108189-65-7D, reaction products with hectorite and montmorillonite

RL: ARU (Analytical role, unclassified); PEP (Physical, engineering or chemical process); TEM (Technical or engineered material use); ANST (Analytical study); PROC (Process); USES (Uses)

(microporous organo-smectites as stationary phases for gas chromatog. separation of alkanes)

RN 108189-65-7 CAPLUS

CN 1,2-Ethanediaminium, N,N,N',N'-tetramethyl-N,N'-bis[2-(trimethylammonio)ethyl]- (9CI) (CA INDEX NAME)



L17 ANSWER 10 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1994:253366 CAPLUS

DOCUMENT NUMBER: 120:253366

TITLE: Compositions and methods for enhanced drug delivery  
Hale, Ron L.; Lu, Amy; Solas, Dennis; Selick, Harold E.; Oldenburg, Kevin R.; Zaffaroni, Alejandro C.

INVENTOR(S): Affymax Technologies N.V., Neth.

PATENT ASSIGNEE(S): PCT Int. Appl., 155 pp.  
CODEN: PIXXD2

DOCUMENT TYPE: Patent  
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9325197	A1	19931223	WO 1993-US5631	19930611 <--
W: AT, AU, BB, BG, BR, CA, CH, CZ, DE, DK, ES, FI, GB, HU, JP, KP, KR, LK, LU, MG, MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SK, UA, US				
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG				
AU 9345345	A1	19940104	AU 1993-45345	19930611 <--
EP 647133	A1	19950412	EP 1993-915319	19930611 <--
R: CH, DE, FR, GB, IT, LI, NL				

US 5607691	A 19970304	US 1995-449188	19950524 <--
PRIORITY APPLN. INFO.:		US 1992-898219	A2 19920612
		US 1993-9463	A2 19930127
		WO 1993-US5631	A 19930611
		US 1993-77296	B2 19930614
		US 1993-164293	B1 19931209

AB The present invention relates to methods of delivering pharmaceutical agents across membranes, including the skin layer or mucosal membranes of a patient. A pharmaceutical agent is covalently bonded to a chemical modifier, via a physiol. cleavable bond, such that the membrane transport and delivery of the agent is enhanced. Progesterone 3-{2-O-[10-O-(O-acetyl-L-carnitiny)decanoyl]glycolic acid} enol ester was prepared from progesterone by preparation of the enol acetate, reaction with 10-hydroxydecanoic acid, and reaction of the hydroxyl diester with 3-O-acetyl-L-carnitine acid chloride (preparation given). In vitro serum half-lives of some pharmaceutical agent-chemical modifier complexes are given.

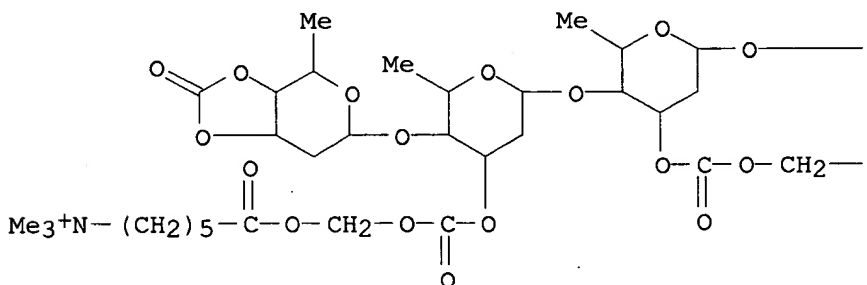
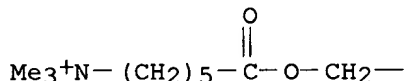
IT 154271-96-2

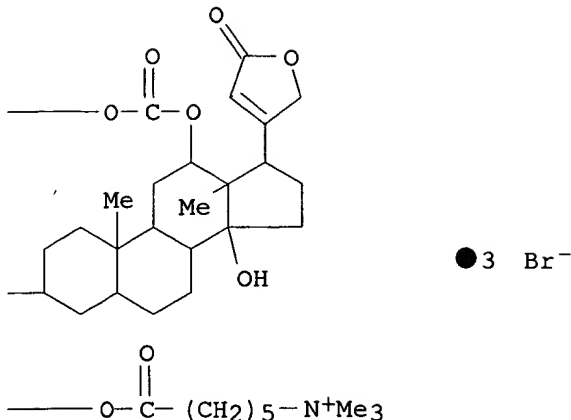
RL: BIOL (Biological study)  
(as drug-chemical modifier conjugate through physiol. cleavable bond, in vitro serum half-life of)

RN 154271-96-2 CAPLUS

CN Card-20(22)-enolide, 3-[[O-3,4-O-carbonyl-2,6-dideoxy- $\beta$ -D-ribo-hexopyranosyl-(1 $\rightarrow$ 4)-O-2,6-dideoxy-3-O-[[[[1-oxo-6-(trimethylammonio)hexyl]oxy]methoxy]carbonyl]- $\beta$ -D-ribo-hexopyranosyl-(1 $\rightarrow$ 4)-2,6-dideoxy-3-O-[[[[1-oxo-6-(trimethylammonio)hexyl]oxy]methoxy]carbonyl]- $\beta$ -D-ribo-hexopyranosyl]oxy]-14-hydroxy-12-[[[[1-oxo-6-(trimethylammonio)hexyl]oxy]methoxy]carbonyl]oxy]-, tribromide, (3 $\beta$ ,5 $\beta$ ,12 $\beta$ )- (9CI) (CA INDEX NAME)

PAGE 1-A





=> d 117 11-20 ibib abs hitstr

L17 ANSWER 11 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1993:131746 CAPLUS  
 DOCUMENT NUMBER: 118:131746  
 TITLE: Shampoos containing cationic and anionic surfactants to impart improved hair conditioning properties  
 INVENTOR(S): Duffy, Michele; Bergmann, Wolfgang  
 PATENT ASSIGNEE(S): Curtis, Helene, Inc., USA  
 SOURCE: Eur. Pat. Appl., 42 pp.  
 CODEN: EPXXDW  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 511652	A1	19921104	EP 1992-107311	19920429 <--
EP 511652	B1	19951129		
R: AT, BE, CH, CA 2066885	DE, DK, ES, FR, AA	19921030	GB, GR, IT, LI, LU, NL, SE CA 1992-2066885	19920423 <--
CA 2066885	C	20020723		
IL 101682	A1	19961205	IL 1992-101682	19920423 <--
NO 9201640	A	19921030	NO 1992-1640	19920428 <--
NO 300355	B1	19970520		
AU 9215224	A1	19921105	AU 1992-15224	19920428 <--
AU 653216	B2	19940922		
ZA 9203084	A	19930127	ZA 1992-3084	19920428 <--
AT 130751	E	19951215	AT 1992-107311	19920429 <--
ES 2080369	T3	19960201	ES 1992-107311	19920429 <--
JP 06107525	A2	19940419	JP 1992-155568	19920430 <--
PRIORITY APPLN. INFO.:			US 1991-692709	A 19910429

OTHER SOURCE(S): MARPAT 118:131746  
 AB A conditioning shampoo comprises (1) an anionic cleansing surfactant 1-15, (2) a polymeric cationic conditioning compound 0.1-2, (3) a cationic conditioning surfactant 0.2-10, (4) a fatty acid ester 0.1-3, and (5) water as carrier. A hair conditioner contained guar hydroxypropyltrimonium 1.50, ricinoleamidopropyl trimonium chloride (Surfactrol Q1) 1.65, linoleamidopropyl PG-dimonium chloride phosphate (Phospholipid EFA) 0.60, ammonium lauryl sulfate 6.14, ammonium lauryl ether sulfate 6.14, cetearyl octanoate (Purcellin oil) 2.00, and water

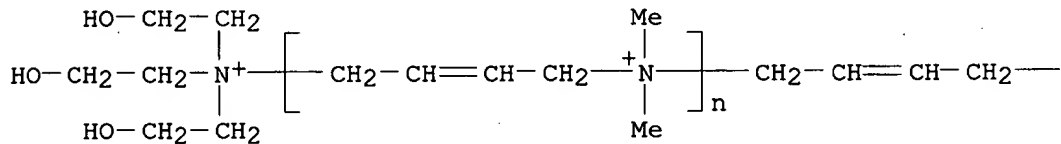
q.s. 100%.

IT 75345-27-6, Polyquaternium 1  
 RL: BIOL (Biological study)  
 (hair conditioning shampoo containing anionic surfactants and fatty acid esters and)

RN 75345-27-6 CAPLUS

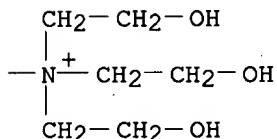
CN Poly[(dimethylimino)-2-butene-1,4-diyl chloride],  $\alpha$ -[4-[tris(2-hydroxyethyl)ammonio]-2-butenyl]- $\omega$ -[tris(2-hydroxyethyl)ammonio]-, dichloride (9CI) (CA INDEX NAME)

PAGE 1-A



● 3 Cl<sup>-</sup>

PAGE 1-B



L17 ANSWER 12 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1992:619747 CAPLUS  
 DOCUMENT NUMBER: 117:219747  
 TITLE: Cosmetic composition containing **quaternary** ammonium functionalized phosphate esters  
 INVENTOR(S): Ziegler, Philip D.; Cheney, Michael C.  
 PATENT ASSIGNEE(S): Chesebrough-Pond's USA Co., USA  
 SOURCE: U.S., 7 pp.  
 CODEN: USXXAM  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5135748	A	19920804	US 1991-662680	19910228 <--
US 5169624	A	19921208	US 1991-662880	19910228 <--
CA 2061679	AA	19920829	CA 1992-2061679	19920221 <--
CA 2061679	C	19970603		
EP 501714	A2	19920902	EP 1992-301511	19920224 <--
EP 501714	A3	19930414		
EP 501714	B1	19970502		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, NL, PT, SE				
AT 152347	E	19970515	AT 1992-301511	19920224 <--
ES 2102458	T3	19970801	ES 1992-301511	19920224 <--
BR 9200637	A	19921110	BR 1992-637	19920226 <--
AU 9211356	A1	19920903	AU 1992-11356	19920228 <--

AU 655229	B2	19941208		
JP 04338312	A2	19921125	JP 1992-43709	19920228 <--
JP 07014848	B4	19950222		
ZA 9201521	A	19930830	ZA 1992-1521	19920228 <--
KR 9701639	B1	19970213	KR 1992-3150	19920228 <--
PRIORITY APPLN. INFO.:			US 1991-662880	19910228
			US 1991-662680	A 19910228

OTHER SOURCE(S): MARPAT 117:219747

AB A cosmetic composition comprises title compds. [RCONH(CH<sub>2</sub>)<sub>3</sub>N+(Me)(Me)CH<sub>2</sub>CH(OH)CH<sub>2</sub>O]P<sub>3</sub>O<sub>3</sub>X- (R = C<sub>5</sub>-17 alkyl, X = anion) 0.10-30, and a cationic polysaccharide 0.1-10%. These compns. are freeze-thaw cycle stable and exhibit unusual skin mildness properties. An emulsion contained cetyl alc. 2.5, glyceryl monostearate 1.5, iso-Pr palmitate 2, petrolatum 2, propylparaben 0.1, water 78.4, glycerin 10, Quatrisoft LM-200 (a cationic polysaccharide 0.25, Monoquat P-TS (phosphate tris alkylamido triquaternary compound) 3, antifoam AF 0.005, methylparaben 0.15, TiO<sub>2</sub> 0.18.

IT 144377-73-1, Phospholipid EFA 144379-29-3, Monoquat P-TS

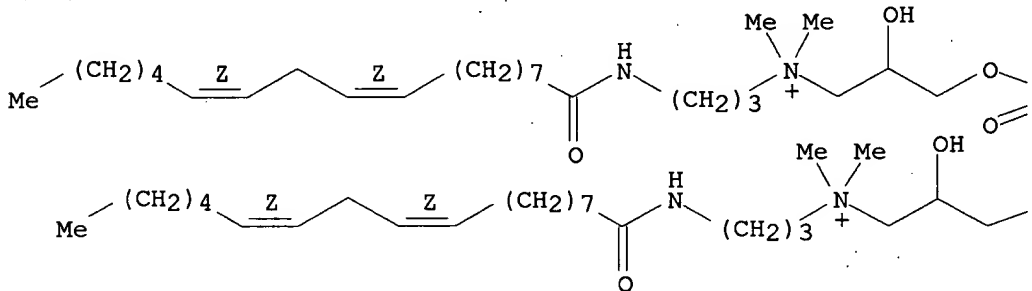
RL: BIOL (Biological study)  
(cosmetic composition containing cationic polysaccharides and)

RN 144377-73-1 CAPLUS

CN 4,6-Dioxa-14-aza-10-azonia-5-phosphadriaconta-23,26-dien-1-aminium, 5-[3-[dimethyl[3-[(9Z,12Z)-1-oxo-9,12-octadecadienyl]amino]propyl]ammonio]-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-15-oxo-N-[3-[(9Z,12Z)-1-oxo-9,12-octadecadienyl]amino]propyl]-, trichloride, 5-oxide, (23Z,26Z)- (9CI) (CA INDEX NAME)

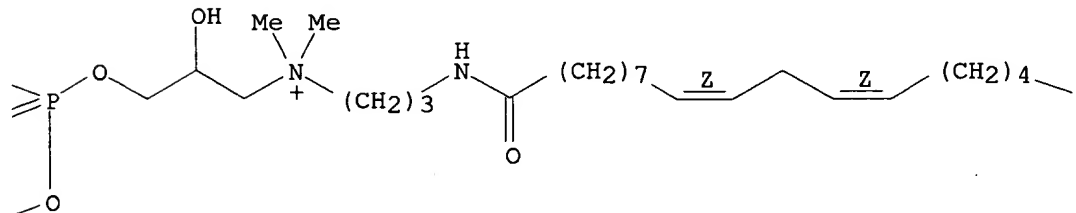
Double bond geometry as shown.

PAGE 1-A



● 3 Cl<sup>-</sup>

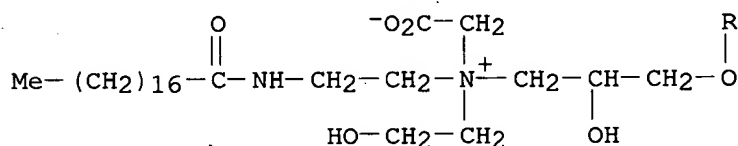
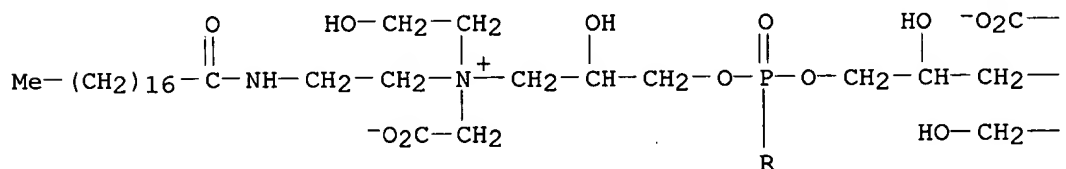
PAGE 1-B



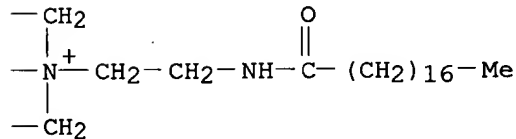
Me

RN 144379-29-3 CAPLUS  
CN 4,6-Dioxa-13-aza-10-azonia-5-phosphahentriacontan-1-aminium,  
N,10-bis(carboxymethyl)-5-[3-[(carboxymethyl)(2-hydroxyethyl)[2-[(1-  
oxooctadecyl)amino]ethyl]ammonio]-2-hydroxypropoxy]-2,8-dihydroxy-N,10-  
bis(2-hydroxyethyl)-14-oxo-N-[2-[(1-oxooctadecyl)amino]ethyl]-, tris(inner  
salt), 5-oxide (9CI) (CA INDEX NAME)

PAGE 1-A



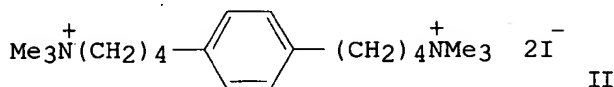
PAGE 1-B



L17 ANSWER 13 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
ACCESSION NUMBER: 1991:514182 CAPLUS  
DOCUMENT NUMBER: 115:114182  
TITLE: Preparation of **quaternary** ammonium compounds  
as muscle relaxants  
INVENTOR(S): Kimura, Masayasu; Naito, Kenji; Sakuma, Osamu; Morita,  
Tadashi  
PATENT ASSIGNEE(S): Tobishi Pharmaceutical Co., Ltd., Japan  
SOURCE: Ger. Offen., 38 pp.  
CODEN: GWXXBX  
DOCUMENT TYPE: Patent  
LANGUAGE: German  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 4010925	A1	19901011	DE 1990-4010925	19900404 <--

JP 02268142	A2	19901101	JP 1989-87889	19890410 <--
CA 2013432	AA	19901010	CA 1990-2013432	19900329 <--
CH 679581	A	19920313	CH 1990-1173	19900405 <--
FR 2645532	A1	19901012	FR 1990-4431	19900406 <--
US 5093370	A	19920303	US 1990-506862	19900409 <--
GB 2230263	A1	19901017	GB 1990-8074	19900410 <--
PRIORITY APPLN. INFO.:			JP 1989-87889	A 19890410
OTHER SOURCE(S):	MARPAT 115:114182			
GI				



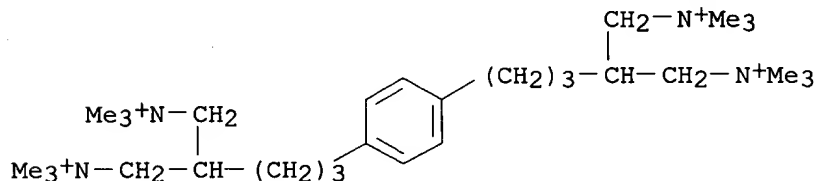
AB [R<sub>2</sub>R<sub>3</sub>QZ(CH<sub>2</sub>)<sub>a</sub>[CH(CH<sub>2</sub>A)CH<sub>2</sub>]bA]<sub>m</sub>+] (R<sub>4</sub>)<sub>m</sub>- [I; Z = CH<sub>2</sub>, alkylene(oxy), alkynylene, CO, CO<sub>2</sub>, alkylene(carbonyloxy), CHOR<sub>5</sub>, alkylene carbonyl, O, S, SO, SO<sub>2</sub>, hydroxyalkyl; R<sub>2</sub> = H, hydroxyalkyl, formyl, alkylcarbonyl, NO<sub>2</sub>, NHR<sub>6</sub>; R<sub>3</sub> = H, Z(CH<sub>2</sub>)<sub>a</sub>[CH(CH<sub>2</sub>A)CH<sub>2</sub>]b; R<sub>4</sub> = anion; R<sub>5</sub>, R<sub>6</sub> = H, Ac; A = **quaternary** ammonium group; Q = trivalent benzene, naphthalene, or biphenyl ring, trivalent ethane radical; a = 1-8; b = 0, 1; m = 1-4] were prepared by reaction of halo derivs. R<sub>2</sub>R<sub>3</sub>QZ(CH<sub>2</sub>)<sub>a</sub>[CH(CH<sub>2</sub>A)CH<sub>2</sub>]bR<sub>7</sub> (R<sub>7</sub> = halo, reactive **ester** group, other symbols as defined above) with tertiary amines. Thus, bromination of 11.5 g 1,4-bis(4-hydroxybutyl)benzene (preparation from p-diiodobenzene given) by PBr<sub>3</sub> gave 13.1 g 1,4-bis(4-bromobutyl)benzene which (3.48 g) was stirred 3.5 h at room temperature with 9 mL 50% Me<sub>2</sub>NH to give 2.70 g 1,4-bis(dimethylamino) analog. This (3.1 g) was refluxed 2.5 h in MeOH with 7.2 g MeI to give 3.57 g title compound II. The latter in vitro inhibited muscle contractions induced by elec. shock with IC<sub>50</sub> of 22.8  $\mu$ M vs. 25.2 and 101  $\mu$ M for succinylcholine and decamethonium, resp. Approx. 42 I were prepared

IT 134519-58-7P

RL: SPN (Synthetic preparation); PREP (Preparation)  
(preparation of, as muscle relaxant)

RN 134519-58-7 CAPLUS

CN 1,4-Benzenedipentanaminium, N,N,N,N',N',N'-hexamethyl- $\beta$ , $\beta'$ -bis[(trimethylammonio)methyl]-, tetraiodide (9CI) (CA INDEX NAME)



● 4 I<sup>-</sup>

L17 ANSWER 14 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1991:209601 CAPLUS

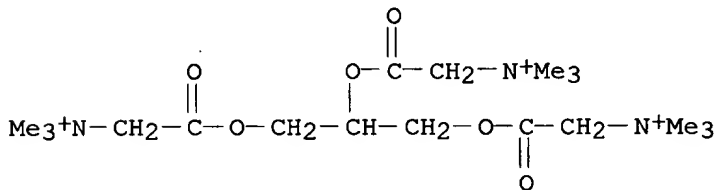
DOCUMENT NUMBER: 114:209601

TITLE: Bleaching detergent compositions containing sulfonate salts

INVENTOR(S): Aoyanagi, Muneo; Kuroda, Mutsumi; Araki, Hiroyuki; Taguchi, Akio

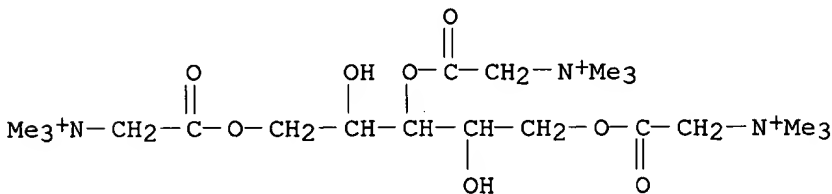
PATENT ASSIGNEE(S): Kao Corp., Japan  
SOURCE: Jpn. Kokai Tokkyo Koho, 9 pp.  
CODEN: JKXXAF  
DOCUMENT TYPE: Patent  
LANGUAGE: Japanese  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 02229895	A2	19900912	JP 1989-230773	19890906 <--
PRIORITY APPLN. INFO.:			JP 1988-237369	A1 19880921
OTHER SOURCE(S):	MARPAT 114:209601			
AB	Title compns. comprise 3-30% peroxides generating H <sub>2</sub> O <sub>2</sub> in aqueous solns., 0.1-30% activators which react with H <sub>2</sub> O <sub>2</sub> to generate cationic group-containing organic peroxy acids, and 10-50% mixts. (9/1-1/3) of alkylbenzenesulfonate salts and $\alpha$ -sulfo fatty acid <b>ester</b> salts. Thus, a composition containing Na dodecylbenzenesulfonate 22, hydrogenated palm oil fatty acid Me <b>ester</b> Na sulfonate 3, Na silicate 5, Na <sub>2</sub> CO <sub>3</sub> 10, 4A zeolite 25, Na <sub>2</sub> S <sub>2</sub> O <sub>8</sub> 10, NCCH <sub>2</sub> N+Me <sub>2</sub> CH <sub>2</sub> CH <sub>2</sub> N+Me <sub>2</sub> CH <sub>2</sub> CN 2Cl <sup>-</sup> 5, PEG 2, protease 2, and H <sub>2</sub> O 5%, with the remainder being Na <sub>2</sub> SO <sub>4</sub> was used to wash and bleach a tea-stained cotton cloth.			
IT	<b>130631-35-5 132787-32-7</b>			
RL:	CAT (Catalyst use); USES (Uses) (activators, for peroxide bleaching agents, in laundry detergents)			
RN	130631-35-5 CAPLUS			
CN	Ethanaminium, 2,2',2'''-[1,2,3-propanetriyltris(oxy)]tris[N,N,N-trimethyl-2-oxo-, trichloride (9CI) (CA INDEX NAME)			



● 3  $\text{Cl}^-$

RN 132787-32-7 CAPLUS  
CN Pentitol, 1,3,5-tris[(trimethylammonio)acetate], trichloride (9CI) (CA  
INDEX NAME)



● 3 Cl<sup>-</sup>

L17 ANSWER 15 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1990:614309 CAPLUS  
 DOCUMENT NUMBER: 113:214309  
 TITLE: Bleaching composition  
 INVENTOR(S): Sotoya, Kohshiro; Ogura, Nobuyuki; Aoyagi, Muneo;  
 Murata, Moriyasu  
 PATENT ASSIGNEE(S): Kao Corp., Japan  
 SOURCE: Eur. Pat. Appl., 18 pp.  
 CODEN: EPXXDW  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 371809	A1	19900606	EP 1989-312492	19891130 <--
EP 371809	B1	19940928		
R: DE, ES, FR, GB				
JP 02147698	A2	19900606	JP 1988-303161	19881130 <--
JP 06096719	B4	19941130		
US 5093022	A	19920303	US 1989-441941	19891127 <--
ES 2059786	T3	19941116	ES 1989-312492	19891130 <--
PRIORITY APPLN. INFO.:			JP 1988-303161	A 19881130

OTHER SOURCE(S): MARPAT 113:214309

AB Compds. R1N+R2R3R4COL X- (R1-3 = alkyl, alkenyl, alkaryl; R4 = alkylene, phenylene, etc.; L = OC6H4CO2R5, OC6H4NR6R7, O-p-C6H4CR6R7-p-C6H4OY, ON:CR5R6, succinimidooxy, etc.; R5 = alkyl; R6-7 = H, alkyl; Y = H, COR4N+R1R2R3; X- = anion) are useful as activators for peroxygen bleaching agents, especially in laundry detergents. The activator Me3N+(CH2)3CO2-p-C6H4CO2Me Cl- was more effective than (Ac2NCH2)2 in the bleaching of tea-stained fabrics with Na percarbonate.

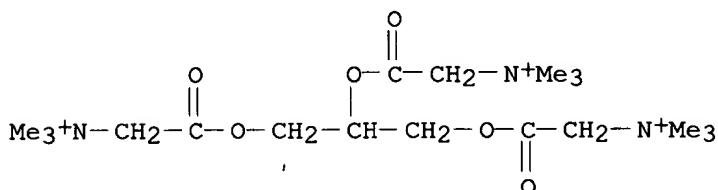
IT 130631-35-5

RL: USES (Uses)

(bleaching activators, for peroxygen compds. in laundrying)

RN 130631-35-5 CAPLUS

CN Ethanaminium, 2,2',2'''-[1,2,3-propanetriyltris(oxy)]tris[N,N,N-trimethyl-2-oxo-, trichloride (9CI) (CA INDEX NAME)



● 3 Cl-

L17 ANSWER 16 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1990:552621 CAPLUS  
 DOCUMENT NUMBER: 113:152621  
 TITLE: Choline esters of alkylenebisphosphonic acids  
 AUTHOR(S): Bikchurina, L. Kh.; Yumagulova, R. Kh.; Khalilov, L. M.; Vasil'eva, E. V.; Leplyanin, G. V.  
 CORPORATE SOURCE: Inst. Khim., Ufa, USSR  
 SOURCE: Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (

1990), (6), 1424-9  
CODEN: IASKA6; ISSN: 0002-3353

DOCUMENT TYPE:

Journal

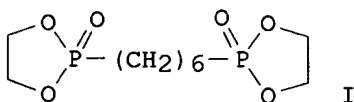
LANGUAGE:

Russian

OTHER SOURCE(S):

CASREACT 113:152621

GI



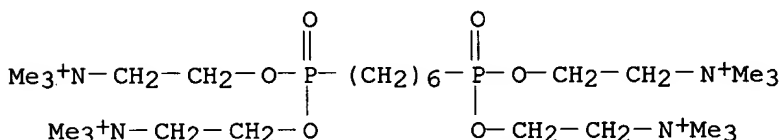
AB Reaction of  $(\text{CH}_2)_6[\text{P}(\text{O})\text{Cl}_2]_2$  with  $\text{HOCH}_2\text{CH}_2\text{OH}$  and  $\text{HOCH}_2\text{CH}_2\text{Cl}$  gave esters I and  $(\text{CH}_2)_6[\text{P}(\text{O})(\text{OCH}_2\text{CH}_2\text{Cl})_2]_2$  resp. which on **quaternization** with  $\text{NMe}_3$  gave  $(\text{CH}_2)_6[\text{P}(\text{O})(\text{O}-)\text{OCH}_2\text{CH}_2\text{N}^+\text{Me}_3]_2$  and  $(\text{CH}_2)_6[\text{P}(\text{O})(\text{OCH}_2\text{N}^+\text{Me}_3\text{Cl}^-)_2]_2$  resp.

IT 129623-00-3P

RL: SPN (Synthetic preparation); PREP (Preparation)  
(preparation of)

RN 129623-00-3 CAPLUS

CN 3,12-Dioxa-4,11-diphosphatetradecane-1,14-diaminium, N,N,N,N',N',N'-hexamethyl-4,11-bis[2-(trimethylammonio)ethoxy]-, tetrachloride, 4,11-dioxide (9CI) (CA INDEX NAME)



● 4 Cl<sup>-</sup>

L17 ANSWER 17 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1988:119376 CAPLUS

DOCUMENT NUMBER: 108:119376

TITLE: Structural variations in amphiphiles: discoidal multivalent cations

AUTHOR(S): Keller-Griffith, R.; Ringsdorf, H.; Vierengel, A.

CORPORATE SOURCE: Inst. Org. Chem., Univ. Mainz, Mainz, D-6500, Fed. Rep. Ger.

SOURCE: Colloid and Polymer Science (1986), 264(11), 924-35

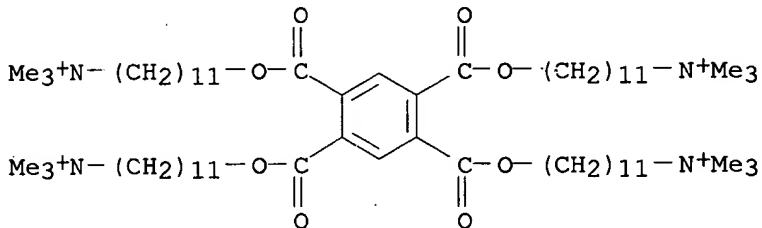
CODEN: CPMSB6; ISSN: 0303-402X

DOCUMENT TYPE: Journal

LANGUAGE: English

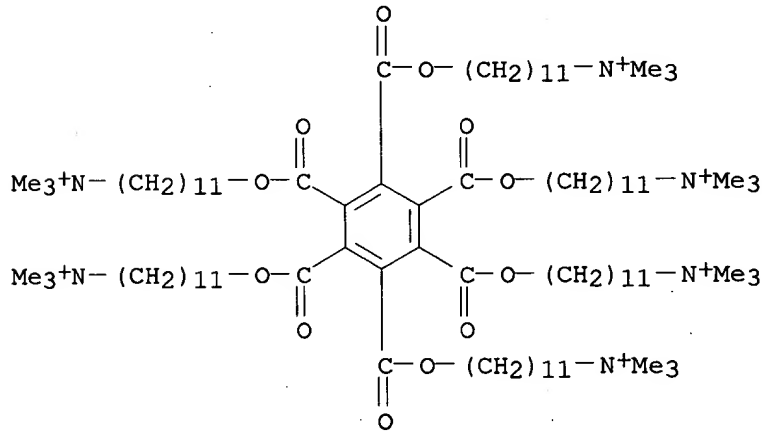
AB Fourteen cationic multipolar amphiphiles were synthesized with pyridinium or trimethylammonium head groups. The hydrophobic cores are planar ring systems (benzene or triphenylene) to which 2, 3, 4, or 6 decylene or undecylene alkyl chains are attached by **ester** linkages. The hydrophilic head groups are bound to the outer ends of the alkyl chains. The aggregation of the mols. in water into micelles and lyotropic liquid crystals was studied. Hexagonal phases are preferred to lamellar phases by these amphiphiles and in more dilute solns. some of these multipolar amphiphiles form cylindrical micelles.

IT 106349-87-5 106349-88-6 113339-63-2  
 113339-65-4 113339-66-5 113339-67-6  
 113339-70-1  
 RL: PRP (Properties)  
 (micelle and liquid crystal aggregation properties of aqueous)  
 RN 106349-87-5 CAPLUS  
 CN 1-Undecanaminium, 11,11',11'',11'''-[1,2,4,5-benzenetetrayltetrakis(carbon yloxy)tetraakis[N,N,N-trimethyl-, tetrabromide (9CI) (CA INDEX NAME)



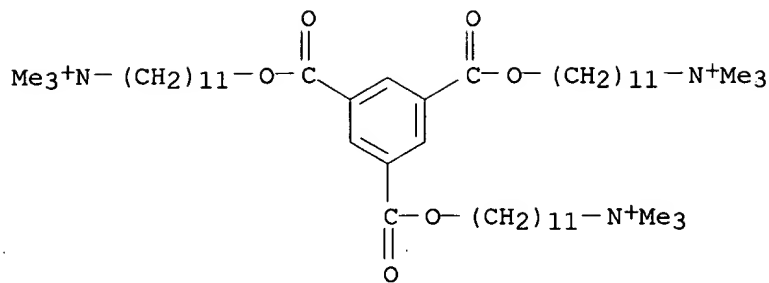
● 4 Br<sup>-</sup>

RN 106349-88-6 CAPLUS  
 CN 1-Undecanaminium, 11,11',11'',11''',11'''',11''''-[1,2,3,4,5,6-benzenehexaylhexakis(carbonyloxy)hexakis[N,N,N-trimethyl-, hexabromide (9CI) (CA INDEX NAME)



● 6 Br<sup>-</sup>

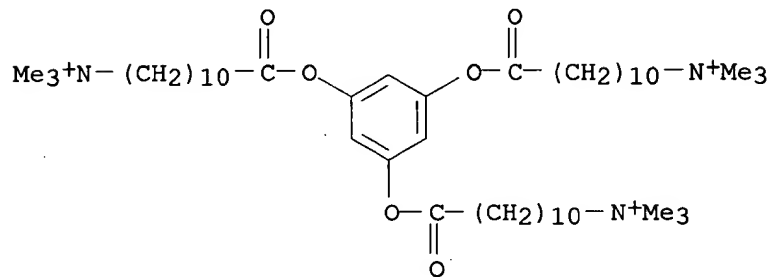
RN 113339-63-2 CAPLUS  
 CN 1-Undecanaminium, 11,11',11''-[1,3,5-benzenetriyltris(carbonyloxy)]tris[N,N,N-trimethyl-, tribromide (9CI) (CA INDEX NAME)



● 3 Br<sup>-</sup>

RN 113339-65-4 CAPLUS

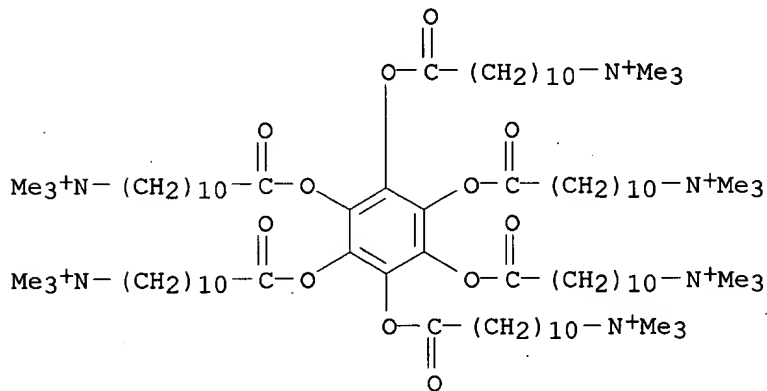
CN 1-Undecanaminium, 11,11',11''-[1,3,5-benzenetriyltris(oxy)]tris[N,N,N-trimethyl-11-oxo-, tribromide (9CI) (CA INDEX NAME)



● 3 Br

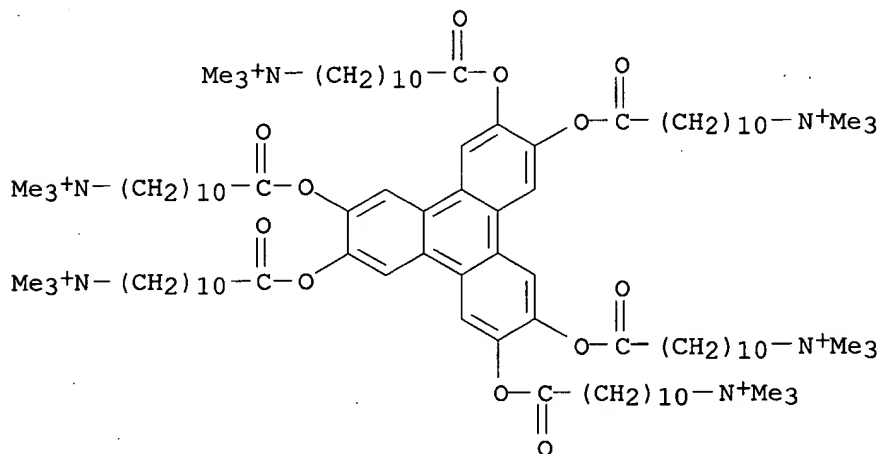
RN 113339-66-5 CAPLUS

CN 1-Undecanaminium, 11,11',11'',11''',11'''',11''''-[1,2,3,4,5,6-benzenehexylhexakis(oxy)]hexakis[N,N,N-trimethyl-11-oxo-, hexabromide (9CI) (CA INDEX NAME)



● 6 Br<sup>-</sup>

RN 113339-67-6 CAPLUS  
 CN 1-Undecanaminium, 11,11',11'',11''',11'''',11''''-[2,3,6,7,10,11-triphenylenehexylhexakis(oxy)]hexakis[N,N,N-trimethyl-11-oxo-, hexabromide (9CI) (CA INDEX NAME)

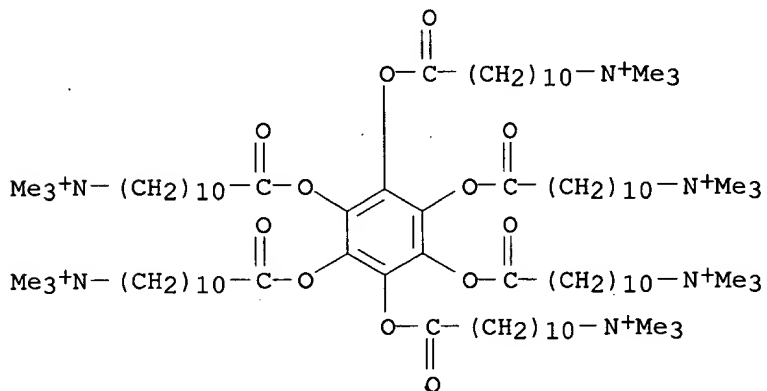


●6 Br<sup>-</sup>

RN 113339-70-1 CAPLUS  
 CN 1-Undecanaminium, 11,11',11'',11''',11'''',11''''-[1,2,3,4,5,6-benzenehexylhexakis(oxy)]hexakis[N,N,N-trimethyl-11-oxo-, salt with 2-hydroxybenzoic acid (1:6) (9CI) (CA INDEX NAME)

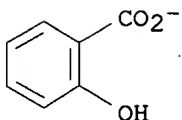
CM 1

CRN 113339-69-8  
 CMF C90 H174 N6 O12



CM 2

CRN 63-36-5  
 CMF C7 H5 O3



L17 ANSWER 18 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1988:75115 CAPLUS

DOCUMENT NUMBER: 108:75115

TITLE: Preparation and formulation of porphyrin derivatives useful for the diagnosis and treatment of cancer

INVENTOR(S): Fukuda, Yozo; Otani, Takuzo; Yamada, Haruo; Sawada, Michikazu; Aizawa, Katsuo; Uchimoto, Mari; Karasawa, Michito

PATENT ASSIGNEE(S): Hamari Chemicals, Ltd., Japan

SOURCE: Eur. Pat. Appl., 33 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

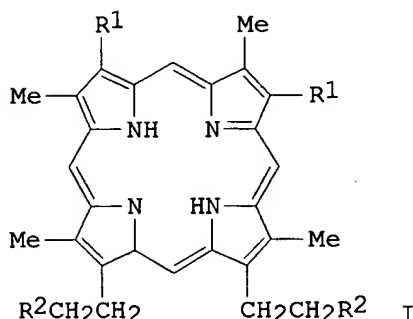
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 233701	A2	19870826	EP 1987-300374	19870116 <--
EP 233701	A3	19880921		
EP 233701	B1	19910814		
R: CH, DE, FR, GB, IT, LI				
JP 62167783	A2	19870724	JP 1986-8789	19860117 <--
JP 07020963	B4	19950308		
JP 62205082	A2	19870909	JP 1986-46000	19860303 <--
JP 07014942	B4	19950222		
JP 63145283	A2	19880617	JP 1986-291904	19861208 <--
PRIORITY APPLN. INFO.:			JP 1986-8789	A 19860117
			JP 1986-46000	A 19860303
			JP 1986-291904	A 19861208

GI



AB Title compds. I [R1 = H, C1-4 alkyl, ethenyl, C2-4Q, Q = di(C1-4-alkyl)amino, tri(C1-4-alkyl)ammonium halide, pyridinio-, quinolinioalkyl halide; R2 = HO2C, C1-4 alkoxy carbonyl, COZ(CmH2m)Q, COZCH(CmH2mQ)2, CH2Q (Z = O, S, HN; m = 1-23]. 7,12-Diethenyl-3,8,13,17-tetramethyl-21H,23H-porphine-2,18-dipropionic acid in CH2Cl2 was treated with (COCl)2 to give the acid chloride which was esterified with Me2NCH2CH2OH to the ester, which in CH2Cl2 was quaternized with MeI to give I [R1 = ethenyl; R2 =

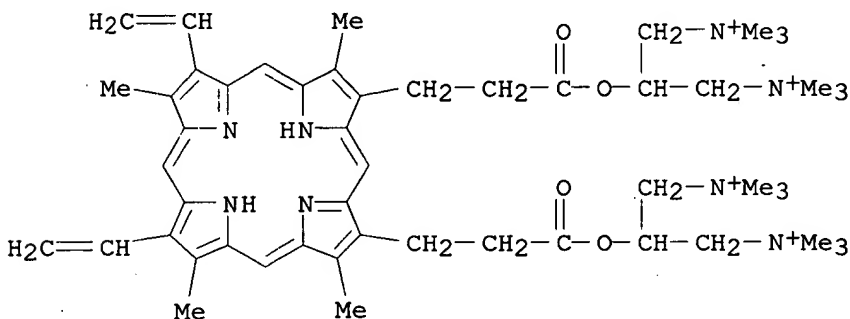
[2-(trimethylammonio)ethoxy]carbonyl diiodide] (II). MKSA cells from mouse nephroadenoma transplanted on a mouse's back, were treated with II at 20 mg/kg, i.v., and excimer laser irradiated, whereby the tumor disappeared after 3 days.

IT 112635-97-9p

RL: SPN (Synthetic preparation); PREP (Preparation)  
(preparation of, as anticancer drug)

RN 112635-97-9 CAPLUS

CN 1,3-Propanediaminium, 2,2'-[ (7,12-diethenyl-3,8,13,17-tetramethyl-21H,23H-porphine-2,18-diyl)bis[(1-oxo-3,1-propanediyl)oxy]]bis[N,N,N,N',N',N'-hexamethyl-, tetraiodide (9CI) (CA INDEX NAME)



●4 I<sup>-</sup>

L17 ANSWER 19 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1987:556318 CAPLUS

DOCUMENT NUMBER: 107:156318

TITLE: Auxiliary agent combination and its use as a textile-finishing agent

INVENTOR(S): Abel, Heinz; Topfl, Rosemarie; Gunter, Franz

PATENT ASSIGNEE(S): Ciba-Geigy A.-G., Switz.

SOURCE: Eur. Pat. Appl., 24 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 225281	A1	19870610	EP 1986-810500	19861103 <--
EP 225281	B1	19890614		
R: BE, CH, DE, FR, GB, IT, LI				
US 4728337	A	19880301	US 1986-925027	19861030 <--
CA 1278402	A1	19910102	CA 1986-522298	19861106 <--
AU 8664951	A1	19870514	AU 1986-64951	19861107 <--
AU 589463	B2	19891012		
ZA 8608485	A	19870624	ZA 1986-8485	19861107 <--
JP 62117887	A2	19870529	JP 1986-264918	19861108 <--
JP 01027189	B4	19890526		

PRIORITY APPLN. INFO.: CH 1985-4802 A 19851108

AB The **quaternary** salts [R<sub>1</sub>COX<sub>1</sub>Z<sub>1</sub>N(R<sub>3</sub>)(R<sub>4</sub>)QN(R<sub>5</sub>)(R<sub>6</sub>)Z<sub>2</sub>X<sub>2</sub>COR<sub>2</sub>]<sup>2+</sup> 2Y<sup>-</sup> (Q = alkylene, optionally containing O or bearing OH; R<sub>1</sub>, R<sub>2</sub> = C<sub>6</sub>-24 aliphatic group; R<sub>3</sub>-6 = alkyl, hydroxyalkyl, alkoxyalkyl; X<sub>1</sub>, X<sub>2</sub> = O, HH; Z<sub>1</sub>, Z<sub>2</sub> = alkylene; Y = anion of a strong acid] are useful in finishing textiles,

especially post-treatment in wool dyeing. Chlorinated wool was dyed with a mixture of chrome, cobalt, and azo dyes, rinsed, heated in an aqueous solution of

0.6% HOCH[CH2N(Me)2(CH2)3NHCOC21H43+]2.2Cl- (I) and 0.6% 4,4'-bis(chloromethyl)biphenyl-N,N,N',N'-tetramethyl-1,6-hexanediamine copolymer (II) at bath ratio 1:30, pH 5, and 40° for 10 min to give a dyeing with fastness to potting, washing, and light 4, 5, and 4-5, resp., and no dry or wet soiling; vs. 4, 5, 4-5, and strong, resp., without I, and 1, 3-4, 4-5, and none, resp., without I and II.

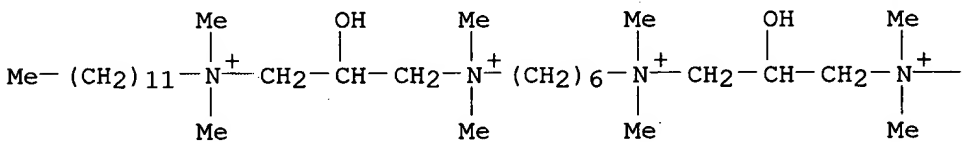
IT 110675-15-5

RL: USES (Uses)  
(afterfinishes, for dyed wool)

RN 110675-15-5 CAPLUS

CN 1,6-Hexanediaminium, N,N'-bis[3-(dodecyldimethylammonio)-2-hydroxypropyl]-N,N,N',N'-tetramethyl-, tetrachloride (9CI) (CA INDEX NAME)

PAGE 1-A



● 4 Cl-

PAGE 1-B

— (CH<sub>2</sub>)<sub>11</sub> — Me

L17 ANSWER 20 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1986:230452 CAPLUS

DOCUMENT NUMBER: 104:230452

TITLE: Antimicrobial compositions for disinfecting surfaces

INVENTOR(S): Gorman, William George; Popp, Karl Frederick

PATENT ASSIGNEE(S): Sterling Drug Inc., USA

SOURCE: Eur. Pat. Appl., 24 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 161425	A2	19851121	EP 1985-103318	19850321 <--
EP 161425	A3	19860226		
R: AT, BE, CH, DE, FR, GB, IT, LI, NL, SE				
AU 8539855	A1	19850926	AU 1985-39855	19850314 <--
JP 60226802	A2	19851112	JP 1985-52175	19850315 <--
ZA 8502023	A	19851127	ZA 1985-2023	19850319 <--
FI 8501139	A	19850924	FI 1985-1139	19850321 <--
NO 8501149	A	19850924	NO 1985-1149	19850321 <--

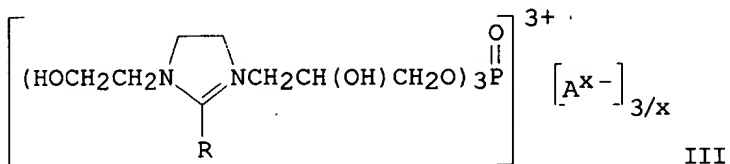
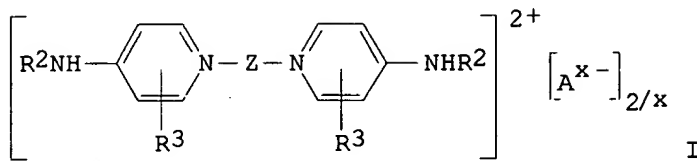
DK 8501321  
IL 74695

A 19850924  
A1 19880630

DK 1985-1321  
IL 1985-74695  
US 1984-592664

19850322 <--  
19850322 <--  
A 19840323

PRIORITY APPLN. INFO.:  
GI



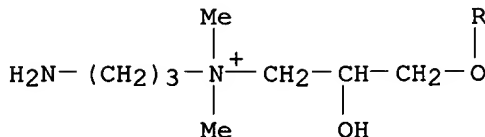
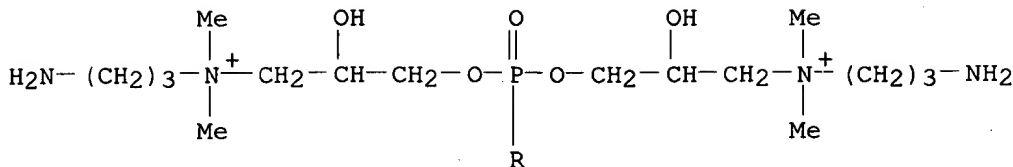
AB The title compns. contain a bisbiguanide, RR<sub>1</sub>NC(:NH)NHC(:NH)NH(CH<sub>2</sub>)<sub>n</sub>NHC(:N H)NHC(:NH)NRR<sub>1</sub> [R = C<sub>6</sub>-16 alkyl, cycloalkyl, polycyclic alkyl, alkylcycloalkyl, cycloalkylalkyl, 4-(2,2-dichlorocyclopropyl)phenyl, (un)substituted Ph; R<sub>1</sub> = H; RR<sub>1</sub> = 3-azabicyclo[3.2.2]nonyl; n = 3-9], or a bis[4-(substituted-amino)-1-pyridinium]alkane I (R<sub>2</sub> = C<sub>6</sub>-18 alkyl, C<sub>5</sub>-7 cycloalkyl, (un)substituted PhCH<sub>2</sub>, Ph; R<sub>3</sub> = H, alkyl; Z = C<sub>4</sub>-18 alkylene; A = anion; x = valence of anion] especially octenidine-HCl, and 1 or more **quaternary** ammonium phosphate **ester** surfactants [(R<sub>4</sub>CONHCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>CH<sub>2</sub>CH(OH)CH<sub>2</sub>O)<sub>3</sub>PO]3+ [Ax-]3/x (II) and III (R<sub>4</sub> = C<sub>5</sub>-17 alkyl; A, x as before) and an aqueous vehicle and addnl. 1 or more polyethylene glycol **ester** surfactant and a **quaternary** nitrogen-containing cellulose ether. Thus, an antimicrobial skin cleansing composition was formulated containing octenidine-HCl 2.0, cocamidopropyl PG-dimonium chloride phosphate (II, R<sub>4</sub>CO = coco acyl) 6.0, PEG-glyceryl cocoate 11.0, NaH<sub>2</sub>PO<sub>4</sub> 0.276, di-Na EDTA 0.1, perfume 0.1, dye 0.005, NaOH to make pH 7.2, and H<sub>2</sub>O to 100% by weight. Porcine skin disks inoculated with *Staphylococcus epidermidis* were immersed in the above composition and the number of surviving bacteria was determined. The results showed a significant mean log<sub>10</sub> count redns. of bacteria on the disks.

IT 75464-22-1D, N-coco acyl derivs.

RL: BIOL (Biological study)  
(skin disinfectant compns. containing surfactants and)

RN 75464-22-1 CAPLUS

CN 4,6-Dioxa-10-azonia-5-phosphatidecan-1-aminium, 13-amino-N-(3-aminopropyl)-5-[3-[(3-aminopropyl)dimethylammonio]-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-, trichloride, 5-oxide (9CI) (CA INDEX NAME)



●3 Cl-

=> d 117 21-30 ibib abs hitstr

L17 ANSWER 21 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1982:533355 CAPLUS

DOCUMENT NUMBER: 97:133355

TITLE: Oily, foaming agent with a liquid phase for care of keratin materials and the skin

INVENTOR(S): Grollier, Jean Francois; Allec, Josiane

PATENT ASSIGNEE(S): Oreal S. A. , Fr.

SOURCE: Ger. Offen., 47 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent

LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 3150338	A1	19820715	DE 1981-3150338	19811218 <--
DE 3150338	C2	19890511		
BE 891528	A1	19820618	BE 1981-206873	19811218 <--
FR 2496458	A1	19820625	FR 1981-23773	19811218 <--
FR 2496458	B1	19870717		
GB 2091100	A	19820728	GB 1981-38210	19811218 <--
GB 2091100	B2	19850227		
JP 57128618	A2	19820810	JP 1981-205048	19811218 <--
JP 02049284	B4	19901029		
CA 1175357	A1	19841002	CA 1981-392613	19811218 <--
US 4488564	A	19841218	US 1981-331904	19811218 <--
CH 651468	A	19850930	CH 1981-8120	19811218 <--
PRIORITY APPLN. INFO.:			LU 1980-83020	A 19801219

AB An oil-containing foaming cleanser for skin and hair contains an oil liquid at ambient temperature 5-85, a surfactant soluble in the oil 15-95, a cationic compound

0.5-10, and H<sub>2</sub>O 0.1-5%. The oil may be plant, animal, or mineral, or synthetic glyceride or fatty acid ester, or fatty alc. The oil-soluble surfactant is anionic, with the acid group neutralized with an amine, or nonionic, and (or) alkanolamide. The cationic compound is a polymer containing polyamino, polyaminoamide, or quaternary ammonium groups as part of the polymer chain. Thus, a shampoo contained: Texapon

WW 99 [83045-95-8] 15, paraffin oil 25, Polymer P1 [68393-49-7] (60% aqueous solution) 3, perfume, antioxidants, and olive oil to 100 g. In use, 20 mL was applied to wet hair, worked in, allowed to stand 10 min, and rinsed to give soft hair that is easily detangled.

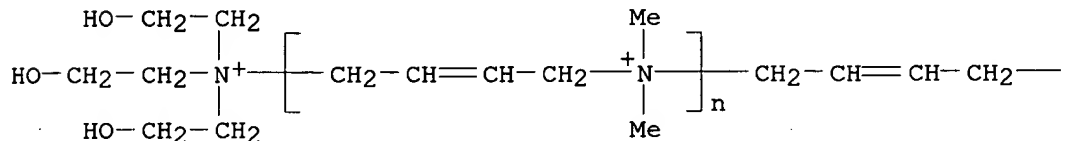
IT 75345-27-6

RL: BIOL (Biological study)  
(shampoos containing oils and)

RN 75345-27-6 CAPLUS

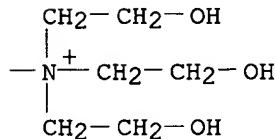
CN Poly[(dimethyliminio)-2-butene-1,4-diyl chloride],  $\alpha$ -[4-[tris(2-hydroxyethyl)ammonio]-2-butenyl]- $\omega$ -[tris(2-hydroxyethyl)ammonio]-, dichloride (9CI) (CA INDEX NAME)

PAGE 1-A



● 3 Cl<sup>-</sup>

PAGE 1-B



L17 ANSWER 22 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1981:426943 CAPLUS

DOCUMENT NUMBER: 95:26943

TITLE: Surfactants useful in conditioning and cleaning agents

INVENTOR(S): Lindeman, Martin K. O.; Stutzman, Ralph; Verdicchio, Robert J.

PATENT ASSIGNEE(S): Johnson and Johnson Baby Products Co., USA

SOURCE: Ger. Offen., 30 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent

LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 3027944	A1	19810219	DE 1980-3027944	19800723 <--
AU 8059761	A1	19810129	AU 1980-59761	19800630 <--
AU 535124	B2	19840301		
FR 2461517	A1	19810206	FR 1980-15681	19800716 <--
CA 1165659	A1	19840417	CA 1980-356656	19800721 <--
GB 2055119	A	19810225	GB 1980-23868	19800722 <--
GB 2055119	B2	19830420		
JP 56020095	A2	19810225	JP 1980-99448	19800722 <--
JP 02025958	B4	19900606		
BR 8004565	A	19810310	BR 1980-4565	19800722 <--

ES 493596	A1	19810616	ES 1980-493596	19800722 <--
ZA 8004422	A	19820224	ZA 1980-4422	19800722 <--
AT 8003794	A	19840215	AT 1980-3794	19800722 <--
AT 375957	B	19840925		

PRIORITY APPLN. INFO.: US 1979-59837 A 19790723

AB **Quaternary** ammonioalkyl phosphates are conditioning agents and detergents in shampoos, wool cleansers, etc., which are nonirritating to skin and eyes. Thus, a shampoo contains OP[OCH<sub>2</sub>CH(OH)CH<sub>2</sub>N+Me<sub>2</sub>C<sub>18</sub>H<sub>37</sub> Cl-]3 [77195-38-1] 2, OP[OCH<sub>2</sub>CH(OH)CH<sub>2</sub>N+Me<sub>2</sub>C<sub>12</sub>H<sub>25</sub> Cl-]3 [77195-39-2] 0.1, C<sub>12</sub>H<sub>25</sub>OSO<sub>3</sub>Na 12, C<sub>11</sub>H<sub>23</sub>CON(CH<sub>2</sub>CH<sub>2</sub>OH)<sub>2</sub> 4, and perfume-dye-water 81.9%.

IT 75464-24-3 77195-35-8 77195-36-9

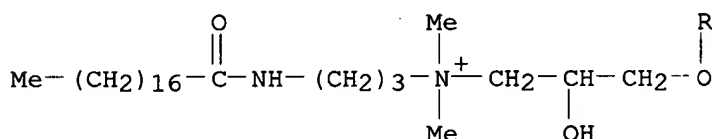
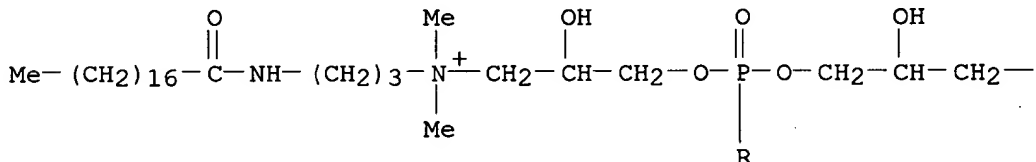
77195-37-0 77195-38-1 77195-39-2

RL: TEM (Technical or engineered material use); USES (Uses)  
(surfactants, nonirritating)

RN 75464-24-3 CAPLUS

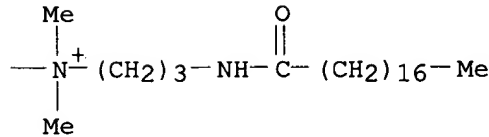
CN 4,6-Dioxa-14-aza-10-azonia-5-phosphadriacontan-1-aminium,  
5-[3-[dimethyl[3-[(1-oxooctadecyl)amino]propyl]ammonio]-2-hydroxypropoxy]-  
2,8-dihydroxy-N,N,10,10-tetramethyl-15-oxo-N-[3-[(1-  
oxooctadecyl)amino]propyl]-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

PAGE 1-A



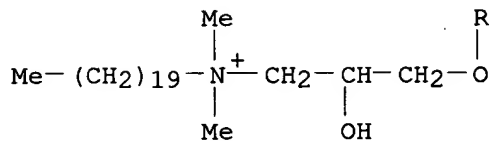
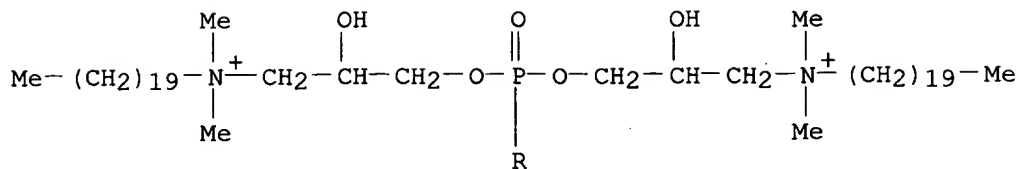
●3 Cl-

PAGE 1-B



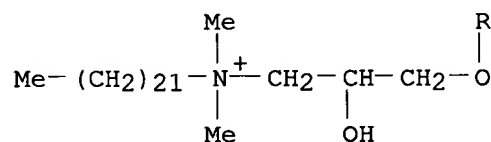
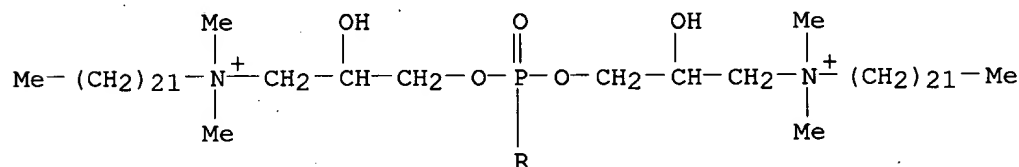
RN 77195-35-8 CAPLUS

CN 4,6-Dioxa-10-azonia-5-phosphadriacontan-1-aminium, N-eicosyl-5-[3-  
(eicosyldimethylammonio)-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-  
tetramethyl-, trichloride, 5-oxide (9CI) (CA INDEX NAME)



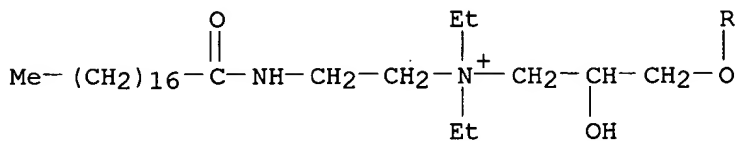
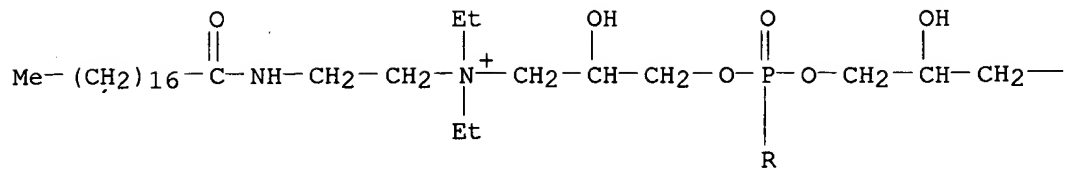
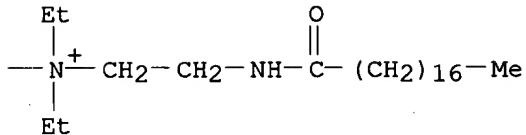
● 3 Cl<sup>-</sup>

RN 77195-36-9 CAPLUS  
 CN 4,6-Dioxa-10-azonia-5-phosphadriacontan-1-aminium, N-docosyl-5-[3-(docosyldimethylammonio)-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-, trichloride, 5-oxide (9CI) (CA INDEX NAME)



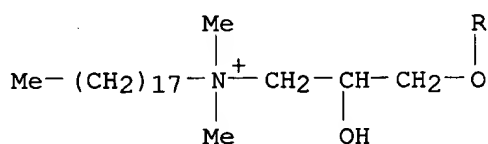
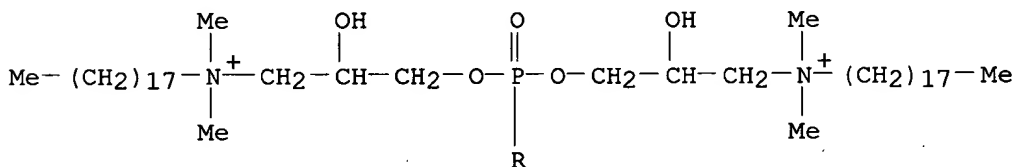
● 3 Cl<sup>-</sup>

RN 77195-37-0 CAPLUS  
 CN 4,6-Dioxa-13-aza-10-azonia-5-phosphadriacontan-1-aminium, 5-[3-[diethyl[2-[(1-oxooctadecyl)amino]ethyl]ammonio]-2-hydroxypropoxy]-N,N,10,10-tetraethyl-2,8-dihydroxy-14-oxo-N-[2-[(1-oxooctadecyl)amino]ethyl]-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

● 3 Cl<sup>-</sup>

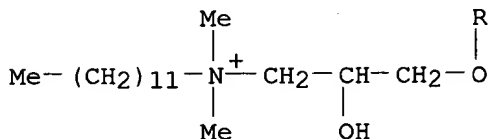
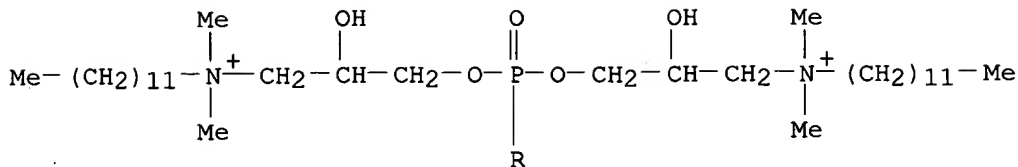
RN 77195-38-1 CAPLUS

CN 4,6-Dioxa-10-azonia-5-phosphaoctacosan-1-aminium, 5-[3-(dimethyloctadecylammonio)-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-N-octadecyl-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

● 3 Cl<sup>-</sup>

RN 77195-39-2 CAPLUS

CN 4,6-Dioxa-10-azonia-5-phosphadocosan-1-aminium, N-dodecyl-5-[3-(dodecyldimethylammonio)-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-, trichloride, 5-oxide (9CI) (CA INDEX NAME)



●3 Cl-

L17 ANSWER 23 OF 43 CAPIPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1981:191667 CAPIPLUS  
 DOCUMENT NUMBER: 94:191667  
 TITLE: Cationic phosphoric acid triesters and their use  
 INVENTOR(S): Lindemann, Martin K. O.; Lukenbach, Elvin R.;  
 Verdicchio, Robert J.  
 PATENT ASSIGNEE(S): Johnson and Johnson Baby Products Co., USA  
 SOURCE: Ger. Offen., 23 pp.  
 CODEN: GWXXBX  
 DOCUMENT TYPE: Patent  
 LANGUAGE: German  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 3027943	A1	19810219	DE 1980-3027943	19800723 <--
DE 3027943	C2	19880901		
AU 8059762	A1	19810129	AU 1980-59762	19800630 <--
AU 536481	B2	19840510		
IN 153525	A	19840721	IN 1980-CA783	19800705 <--
FR 2461715	A1	19810206	FR 1980-15682	19800716 <--
CA 1126750	A1	19820629	CA 1980-356655	19800721 <--
BR 8004564	A	19810203	BR 1980-4564	19800722 <--
JP 56022791	A2	19810303	JP 1980-99449	19800722 <--
JP 01008634	B4	19890214		
ES 493597	A1	19810616	ES 1980-493597	19800722 <--
ZA 8004419	A	19820224	ZA 1980-4419	19800722 <--

PRIORITY APPLN. INFO.: US 1979-59838 A 19790723

AB Title compds. were prepared for use in hair preps. Thus, 19.0 g 85.5% H<sub>3</sub>PO<sub>4</sub> were added dropwise to 46.53 g epichlorohydrin at 80-5°, the mixture was heated 1 h at 80°, 148 g C<sub>18</sub>H<sub>37</sub>NMe<sub>2</sub> were added, and the mixture was heated 20 h at 102° to give OP[OCH<sub>2</sub>CH(OH)CH<sub>2</sub>NMe<sub>2</sub>C<sub>18</sub>H<sub>37</sub>]Cl 3, 98.6% pure.

IT 75464-24-3P 77195-37-0P 77195-38-1P  
 77195-39-2P 77583-79-0P 77583-80-3P

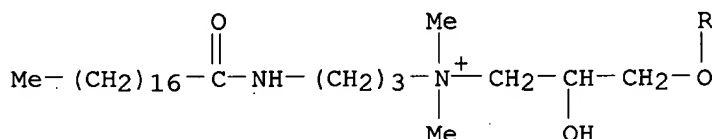
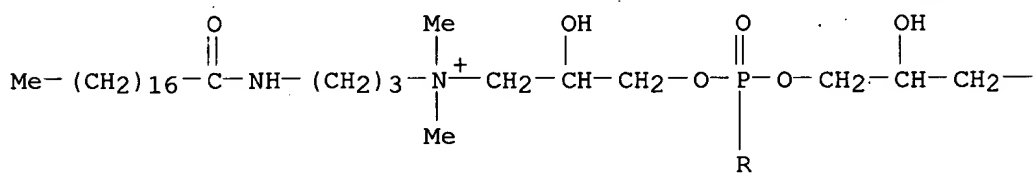
77593-31-8P  
 RL: SPN (Synthetic preparation); PREP (Preparation)  
 (manufacture of, for use in hair preps.)

RN 75464-24-3 CAPIPLUS

CN 4,6-Dioxa-14-aza-10-azonia-5-phosphadotriacontan-1-aminium,

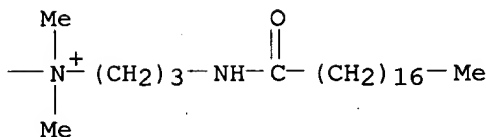
5-[3-[dimethyl[3-[(1-oxooctadecyl)amino]propyl]ammonio]-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-15-oxo-N-[3-[(1-oxooctadecyl)amino]propyl]-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

PAGE 1-A



●3 Cl<sup>-</sup>

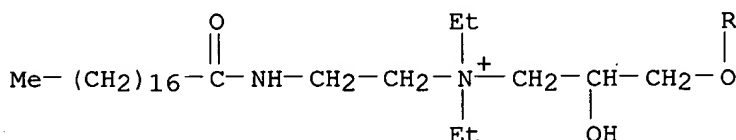
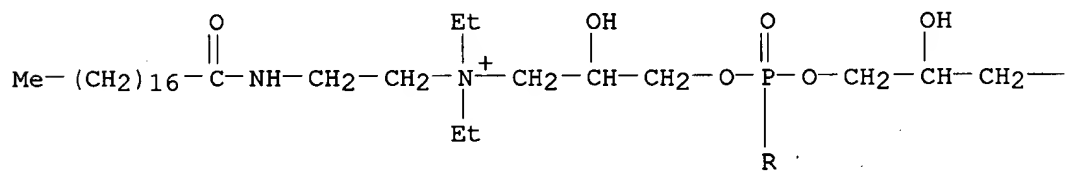
PAGE 1-B



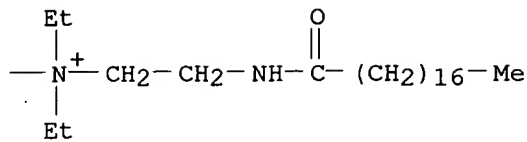
RN 77195-37-0 CAPLUS

CN 4,6-Dioxa-13-aza-10-azonia-5-phosphahentriacontan-1-aminium,  
5-[3-[diethyl[2-[(1-oxooctadecyl)amino]ethyl]ammonio]-2-hydroxypropoxy]-  
N,N,10,10-tetraethyl-2,8-dihydroxy-14-oxo-N-[2-[(1-  
oxooctadecyl)amino]ethyl]-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

PAGE 1-A

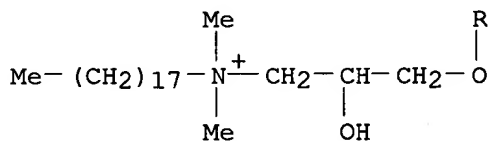
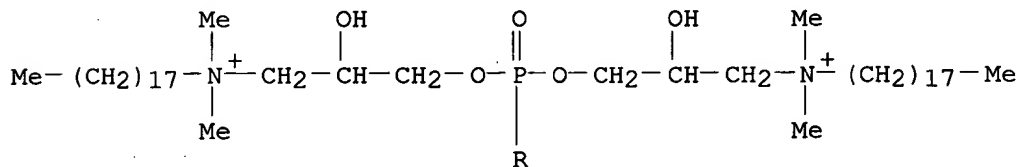


●3 Cl<sup>-</sup>



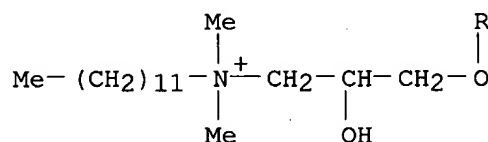
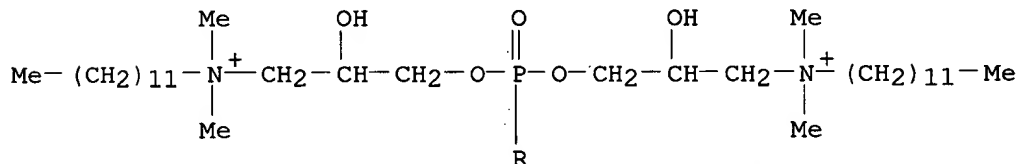
RN 77195-38-1 CAPLUS

CN 4,6-Dioxa-10-azonia-5-phosphaoctacosan-1-aminium, 5-[3-(dimethyloctadecylammonio)-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-N-octadecyl-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

● 3 Cl<sup>-</sup>

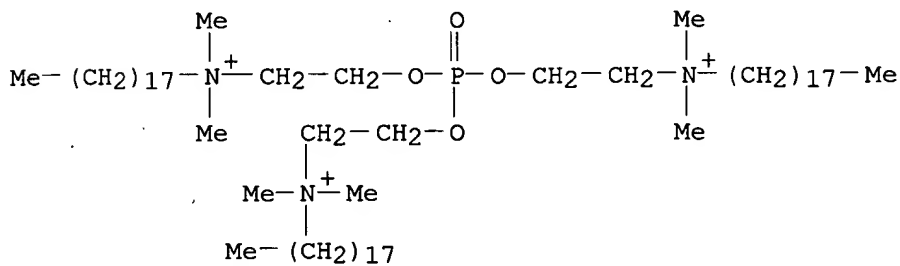
RN 77195-39-2 CAPLUS

CN 4,6-Dioxa-10-azonia-5-phosphadocosan-1-aminium, N-dodecyl-5-[3-(dodecyldimethylammonio)-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

● 3, Cl<sup>-</sup>

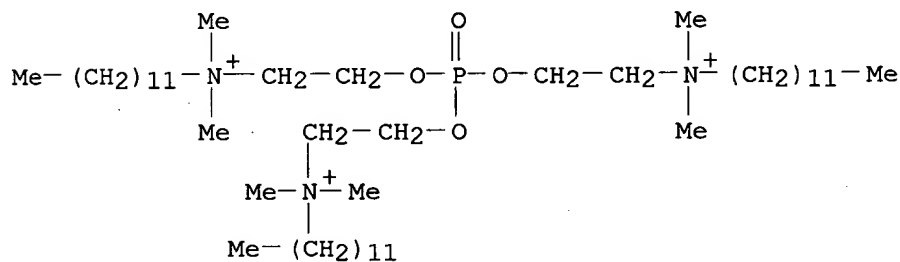
RN 77583-79-0 CAPLUS

CN 3,5-Dioxa-8-azonia-4-phosphahexacosan-1-aminium, 4-[2-(dimethyloctadecylammonio)ethoxy]-N,N,8,8-tetramethyl-N-octadecyl-, trichloride, 4-oxide (9CI) (CA INDEX NAME)



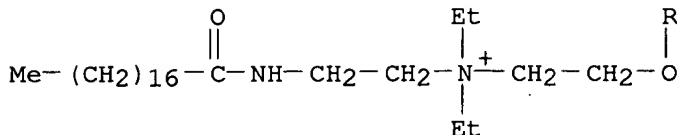
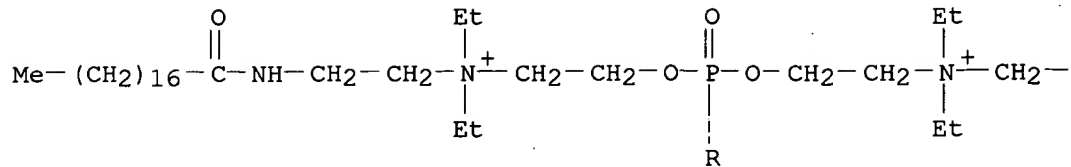
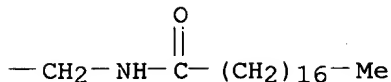
• 3 Cl<sup>-</sup>

RN 77583-80-3 CAPIUS  
CN 3,5-Dioxa-8-azonia-4-phosphaeicosan-1-aminium, N-dodecyl-4-[2-(dodecyldimethylammonio)ethoxy]-N,N,8,8-tetramethyl-, trichloride, 4-oxide (9CI) (CA INDEX NAME)



• 3 Cl<sup>-</sup>

RN 77593-31-8 CAPLUS  
CN 3,5-Dioxa-11-aza-8-azonia-4-phosphonacosan-1-aminium,  
4-[2-[diethyl[2-[(1-oxooctadecyl)amino]ethyl]ammonio]ethoxy]-N,N,8,8-  
tetraethyl-12-oxo-N-[2-[(1-oxooctadecyl)amino]ethyl]-, trichloride,  
4-oxide (9CI) (CA INDEX NAME)

● 3 Cl<sup>-</sup>

L17 ANSWER 24 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1980:604049 CAPLUS

DOCUMENT NUMBER: 93:204049

TITLE: Phosphate **quaternary** compounds

INVENTOR(S): Mayhew, Raymond L.; O'Lenick, Anthony J.

PATENT ASSIGNEE(S): Mona Industries, Inc., USA

SOURCE: U.S., 6 pp.

CODEN: USXXAM

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4209449	-----	19800624	US 1978-965458	19781130 <--

AB The **quaternary** compds. [R3N+CH2CH(OH)CH2O]3PO.3X- (R = alkyl, substituted alkyl, amidoalkyl; R3N = heterocyclic moiety; X = e.g., Cl), useful in detergents, cosmetics, wetting agents, etc., were prepared. Thus, stirring [ClCH2CH(OH)CH2O]3PO with RCONH(CH2)3NMe2 (RCONH = cocamido) gave [RCONH(CH2)3N+Me2CH2CH(OH)CH2O]3PO.3Cl-.

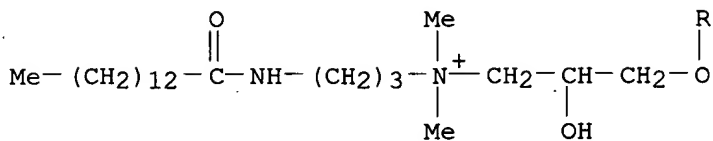
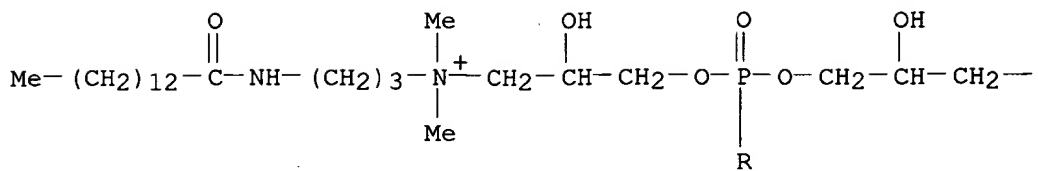
IT 75464-23-2P 75464-24-3P

RL: SPN (Synthetic preparation); PREP (Preparation)  
(preparation and surfactant properties of)

RN 75464-23-2 CAPLUS

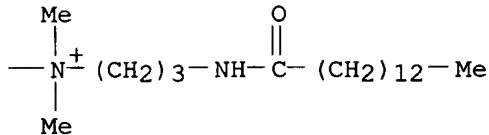
CN 4,6-Dioxa-14-aza-10-azonia-5-phosphaoctacosan-1-aminium,  
5-[3-[dimethyl[3-[(1-oxotetradecyl)amino]propyl]ammonio]-2-hydroxypropoxy]-  
2,8-dihydroxy-N,N,10,10-tetramethyl-15-oxo-N-[3-[(1-  
oxotetradecyl)amino]propyl]-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

PAGE 1-A



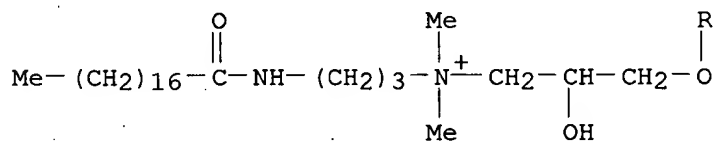
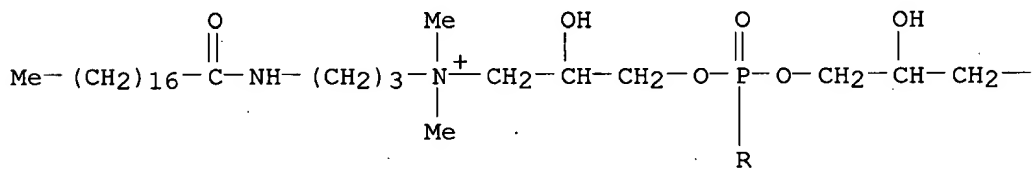
● 3 Cl<sup>-</sup>

PAGE 1-B

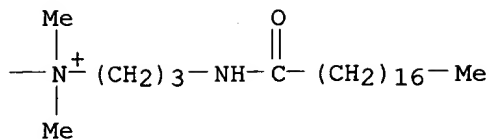


RN 75464-24-3 CAPLUS  
CN 4,6-Dioxa-14-aza-10-azonia-5-phosphadriacontan-1-aminium,  
5-[3-[dimethyl[3-[(1-oxooctadecyl)amino]propyl]ammonio]-2-hydroxypropoxy]-  
2,8-dihydroxy-N,N,10,10-tetramethyl-15-oxo-N-[3-[(1-  
oxooctadecyl)amino]propyl]-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

PAGE 1-A



● 3 Cl<sup>-</sup>



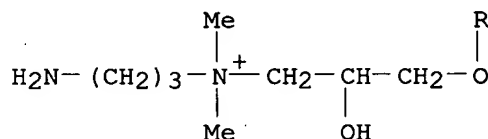
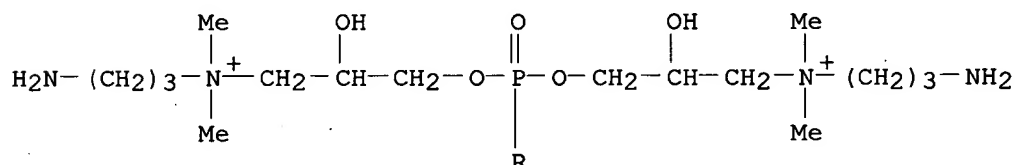
IT 75464-22-1DP, N-cocoyl derivs. 75464-26-5P

75464-27-6P 75477-65-5P 77195-39-2P

RL: SPN (Synthetic preparation); PREP (Preparation)  
(preparation of)

RN 75464-22-1 CAPLUS

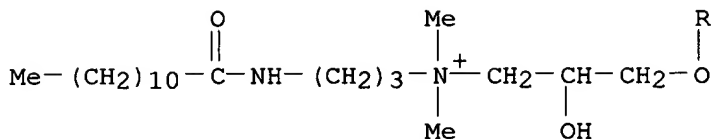
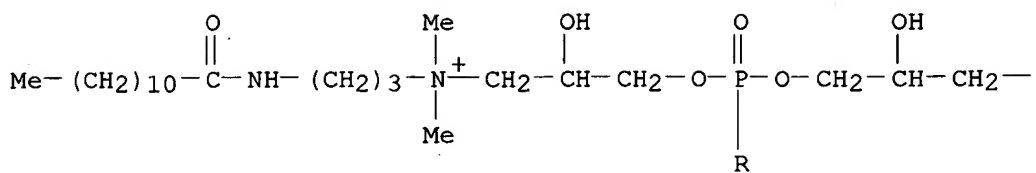
CN 4,6-Dioxa-10-azonia-5-phosphatidecan-1-aminium, 13-amino-N-(3-aminopropyl)-5-[3-[(3-aminopropyl)dimethylammonio]-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

●3 Cl<sup>-</sup>

RN 75464-26-5 CAPLUS

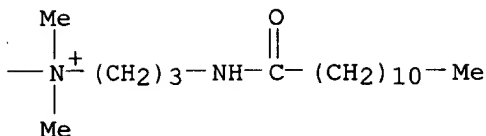
CN 4,6-Dioxa-10-azonia-5-phosphatetraacosan-1-aminium, 5-[3-(dimethyltetradecylammonio)-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-N-tetradecyl-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

PAGE 1-A



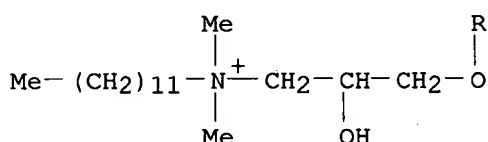
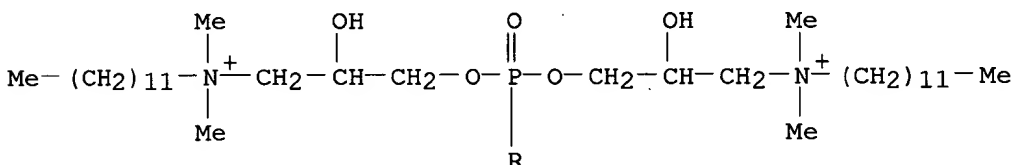
● 3 Cl<sup>-</sup>

PAGE 1-B



RN 77195-39-2 CAPLUS

CN 4,6-Dioxa-10-azonia-5-phosphadocosan-1-aminium, N-dodecyl-5-[3-(dodecyldimethylammonio)-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-, trichloride, 5-oxide (9CI) (CA INDEX NAME)



● 3 Cl<sup>-</sup>

L17 ANSWER 25 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

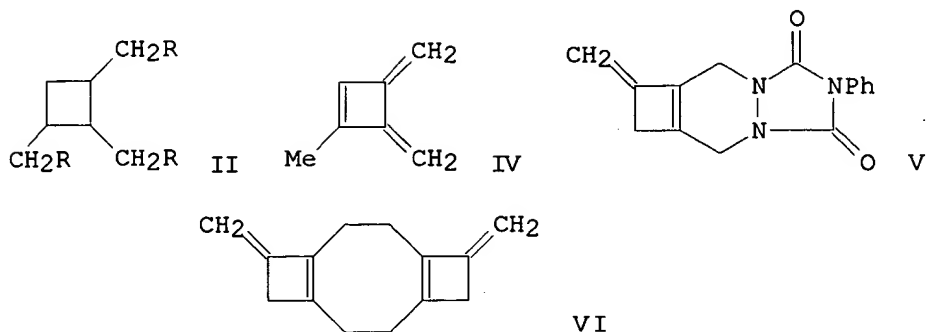
ACCESSION NUMBER: 1980:75908 CAPLUS

DOCUMENT NUMBER: 92:75908

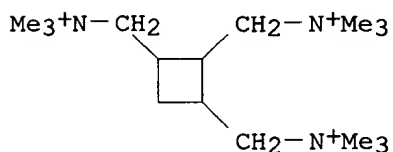
TITLE: Small rings. Part 31. Trimethylenecyclobutane

AUTHOR(S): Martin, Hans Dieter; Mayer, Bernhard

CORPORATE SOURCE: Inst. Org. Chem., Univ. Wuerzburg, Wuerzburg, D-8700,  
 Fed. Rep. Ger.  
 SOURCE: Tetrahedron Letters (1979), (25), 2351-2  
 DOCUMENT TYPE: Journal  
 LANGUAGE: German  
 OTHER SOURCE(S): CASREACT 92:75908  
 GI



AB The title compound (I) was prepared (60%) in 7 steps from  $\text{EtO}_2\text{CCH}_2\text{CH}_2\text{CBr}_2\text{CO}_2\text{Et}$ , the key step being Cope elimination of the cyclobutane derivative II [ $\text{R} = \text{N}(\text{O})\text{Me}_2$ ] (III). Quaternization of III with  $\text{MeI}$  gave II ( $\text{R} = \text{N}^+\text{Me}_3\text{OH}^-$ ), which on thermolysis ( $80-130^\circ$ ) gave the cyclobutene derivative IV. I reacted with N-phenyl-1,2,4-triazoline-3,5-dione to give V and dimerized to give VI.  
 IT 72672-10-7P  
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)  
 (preparation and thermolysis of)  
 RN 72672-10-7 CAPLUS  
 CN 1,2,3-Cyclobutanetrimethanaminium, N,N,N,N',N',N'',N'',N'''-nonamethyl-, trihydroxide (9CI) (CA INDEX NAME)



●3 OH<sup>-</sup>

L17 ANSWER 26 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1977:585212 CAPLUS  
 DOCUMENT NUMBER: 87:185212  
 TITLE: Catalyses by polymer complexes. V. The heterotropic (allosteric) interaction of histamine- and hydroxamate-containing polymer catalysts with hydrophobic ammonium salts in the hydrolysis of phenyl esters  
 AUTHOR(S): Shinkai, Seiji; Tou, Kunio; Kunitake, Toyoki

CORPORATE SOURCE: Fac. Eng., Kyushu Univ., Fukuoka, Japan  
 SOURCE: Polymer Journal (Tokyo, Japan) (1977), 9(4),  
 381-9  
 CODEN: POLJB8; ISSN: 0032-3896  
 DOCUMENT TYPE: Journal  
 LANGUAGE: English  
 AB Methacrylic acid-N-methacryloylhistamine copolymer (I) [64541-47-5],  
 methacrylamide-N-methacryloylhistamine copolymer [64541-48-6] and  
 methacrylic acid-methacrylohydroxamic acid copolymer (II) [64541-49-7]  
 were prepared and their hydrolytic reactivities toward p-nitrophenyl acetate  
 (III) [830-03-5] and p-nitrophenyl hexanoate (IV) [956-75-2] were studied  
 in the absence and in the presence of hydrophobic ammonium salts. The  
 nucleophilic reactivity of I toward III was hardly affected by the addition  
 of hydrophobic ammonium ions, while a marked increase of rates was found  
 in the reaction with IV. Addition of these ammonium ions to the II system  
 enhanced the rate of reaction with IV by lowering the pKa (0.3-0.4 pK  
 unit) and by increasing the second-order rate constant (.apprx.3-fold), as  
 inferred from the pH-rate profile. The rate-enhancing effect of the  
 hydrophobic ammonium salts was analyzed by using the Hill equation which  
 has been employed for analyzing allosteric behavior in enzyme systems; the  
 observed coefficient (n = 3-4) suggested that polymer-bound ammonium ions  
 facilitated the subsequent binding.

IT 64554-59-2 64554-60-5 64554-61-6

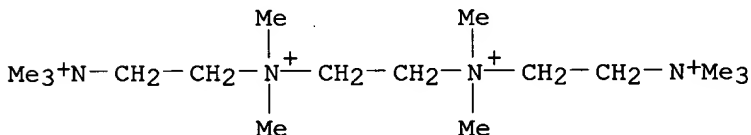
64596-41-4 64596-42-5

RL: PRP (Properties)

(heterotropic interaction of, with histamine- and hydroxamate-containing  
 polymer catalysts, in hydrolysis of phenyl esters)

RN 64554-59-2 CAPLUS

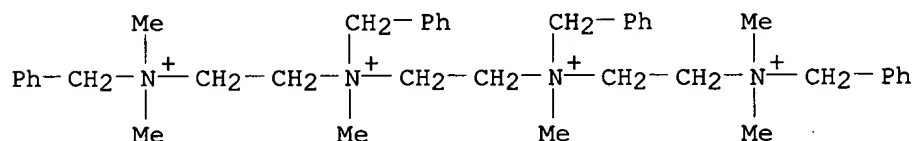
CN 1,2-Ethanediuminium, N,N,N',N'-tetramethyl-N,N'-bis[2-  
 (trimethylammonio)ethyl]-, tetrabromide (9CI) (CA INDEX NAME)



● 4 Br<sup>-</sup>

RN 64554-60-5 CAPLUS

CN 1,2-Ethanediuminium, N,N'-bis[2-[dimethyl(phenylmethyl)ammonio]-N,N'-  
 dimethyl-N,N'-bis(phenylmethyl)-, tetrabromide (9CI) (CA INDEX NAME)

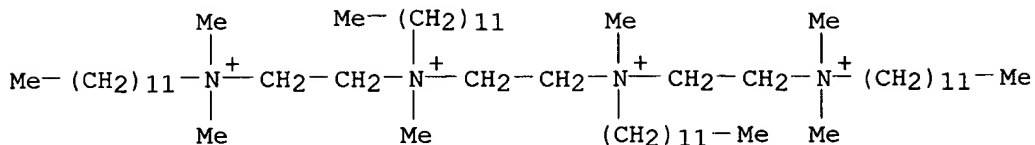


● 4 Br<sup>-</sup>

RN 64554-61-6 CAPLUS

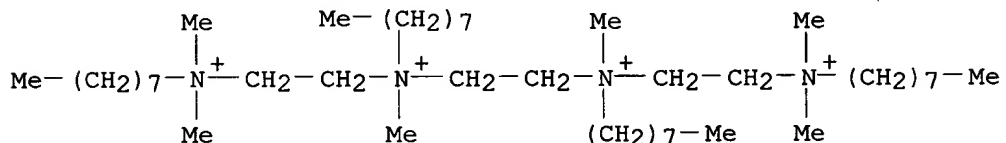
CN 1,2-Ethanediuminium, N,N'-didodecyl-N,N'-bis[2-  
 (dodecyldimethylammonio)ethyl]-N,N'-dimethyl-, tetrabromide (9CI) (CA

INDEX NAME)



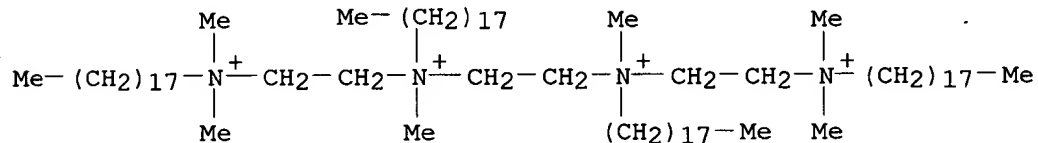
● 4 Br<sup>-</sup>

RN 64596-41-4 CAPLUS  
CN 1,2-Ethanediinium, N,N'-bis[2-(dimethyloctylammonio)ethyl]-N,N'-dimethyl-N,N'-dioctyl-, tetrabromide (9CI) (CA INDEX NAME)



● 4 Br<sup>-</sup>

RN 64596-42-5 CAPLUS  
CN 1,2-Ethanediinium, N,N'-bis[2-(dimethyloctadecylammonio)ethyl]-N,N'-dimethyl-N,N'-dioctadecyl-, tetrabromide (9CI) (CA INDEX NAME)



● 4 Br<sup>-</sup>

L17 ANSWER 27 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1975:589843 CAPLUS

DOCUMENT NUMBER: 83:189843

TITLE: Reactivation and aging of diphenyl phosphoryl acetylcholinesterase

AUTHOR(S): Maglothin, James A.; Wins, Pierre; Wilson, Irwin B.

CORPORATE SOURCE: Dep. Chem., Univ. Colorado, Boulder, CO, USA

SOURCE: Biochimica et Biophysica Acta (1975), 403(2), 370-87

CODEN: BBACAQ; ISSN: 0006-3002

DOCUMENT TYPE: Journal

LANGUAGE: English

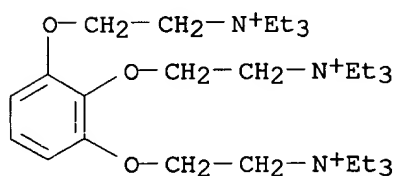
AB Acetylcholinesterase (EC 3.1.1.7) was readily inhibited by 10-5M diphenylphosphorochloridate even though the inhibitor hydrolyzes in a few seconds. The fluoridate was a much weaker inhibitor. The inhibited enzyme, diphenylphosphoryl enzyme spontaneously recovered only .apprx.50% of its activity with a half time of .apprx.17 min at pH 7.0 and 6 min at pH 8.0. The fact that only 50% of the original activity returns was due to aging. The rates of reactivation and aging were very greatly increased by a few percent of an organic solvent. Depending on the solvent, even 1% may increase the rates by a factor of 5-6. The highest increase in rate was 70-fold. **Quaternary** NH4+ also increased the rates. Organic solvents and NH4+ also accelerated the reactivation of the much more stable diethylphosphoryl enzyme derivative

IT 65-29-2

RL: BIOL (Biological study)  
(aging and reactivation of diphenylphosphoryl acetylcholinesterase response to)

RN 65-29-2 CAPLUS

CN Ethanaminium, 2,2',2'''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-, triiodide (9CI) (CA INDEX NAME)



●3 I-

L17 ANSWER 28 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1964:34024 CAPLUS

DOCUMENT NUMBER: 60:34024

ORIGINAL REFERENCE NO.: 60:6087d-e

TITLE: The excitation of lateral geniculate neurons by **quaternary** ammonium derivatives

AUTHOR(S): Curtis, D. R.; Davis, R.

CORPORATE SOURCE: Australian Natl. Univ., Canberra

SOURCE: Journal of Physiology (Cambridge, United Kingdom) (1963), 165(1), 62-82

CODEN: JPHYA7; ISSN: 0022-3751

DOCUMENT TYPE: Journal

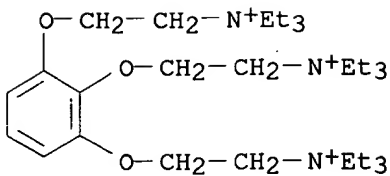
LANGUAGE: Unavailable

AB Carbamoylcholine was the most active excitant tested on cat neurons. Synaptic excitation by the optic nerve, but not by acetylcholine, was suppressed by 5-hydroxytryptamine; dihydro-β-erythroidine had the inverse effect.

IT 65-29-2, [v-Phenyltris(oxyethylene)]tris[triethylammonium iodide]  
(nerve response to)

RN 65-29-2 CAPLUS

CN Ethanaminium, 2,2',2'''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-, triiodide (9CI) (CA INDEX NAME)



●3 I-

L17 ANSWER 29 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1963:434890 CAPLUS

DOCUMENT NUMBER: 59:34890

ORIGINAL REFERENCE NO.: 59:6196e-f

TITLE: The use of paper chromatographic methods for the toxicological determination of drugs. III. Paper chromatographic behavior of several basic drugs as affected by their structure

AUTHOR(S): Vecerkova, J.; Solc, J.; Kacl, K.

CORPORATE SOURCE: Karlova Univ., Prague

SOURCE: Journal of Chromatography (1963), 10, 479-92

CODEN: JOCRAM; ISSN: 0021-9673

DOCUMENT TYPE: Journal

LANGUAGE: German

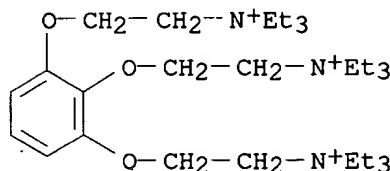
AB cf. CA 58, 2326b, 3714f. The relation between the structure and paper chromatographic behavior of 30 basic drugs was studied; using the reverse phase system petroleum (b.p. 195-220°)-EtOH-H<sub>2</sub>ONH<sub>3</sub>, in which the proportions of EtOH and H<sub>2</sub>O were varied 7 times. In all cases except 3, R<sub>f</sub> increased with increasing EtOH content for 20 tertiary bases and decreased for 10 **quaternary** bases. For most compds., the R<sub>f</sub> was lower at 12-13° than at 17°. R<sub>f</sub> values in the 7 solvent systems are tabulated for all compds.

IT 65-29-2, [v-Phenyltris(oxyethylene)]tris[triethylammonium iodide]

(chromatography of)

RN 65-29-2 CAPLUS

CN Ethanaminium, 2,2',2''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-, triiodide (9CI) (CA INDEX NAME)



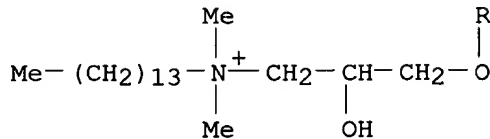
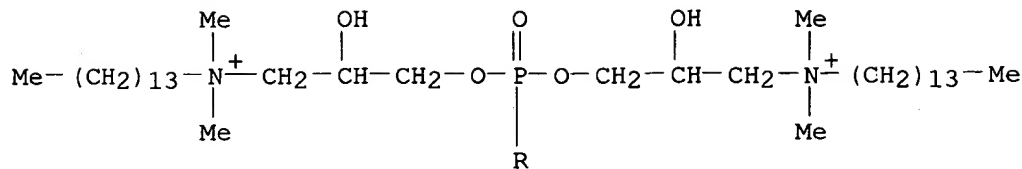
●3 I-

L17 ANSWER 30 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1963:415185 CAPLUS

DOCUMENT NUMBER: 59:15185

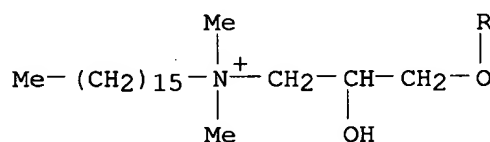
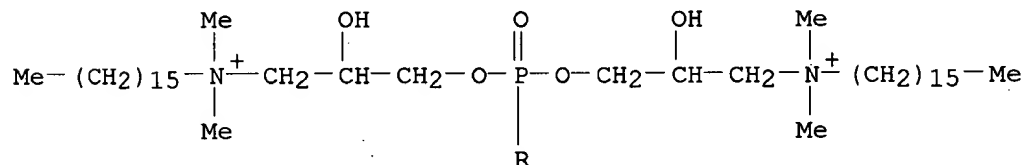
ORIGINAL REFERENCE NO.: 59:2657c-d



● 3 Cl<sup>-</sup>

RN 75464-27-6 CAPLUS

CN 4,6-Dioxa-10-azonia-5-phosphahexacosan-1-aminium, N-hexadecyl-5-[3-(hexadecyldimethylammonio)-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

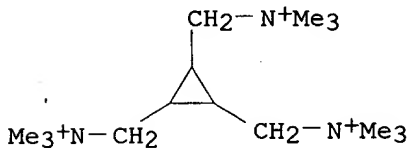


● 3 Cl<sup>-</sup>

RN 75477-65-5 CAPLUS

CN 4,6-Dioxa-14-aza-10-azonia-5-phosphahexacosan-1-aminium, 5-[3-[dimethyl[3-[(1-oxododecyl)amino]propyl]ammonio]-2-hydroxypropoxy]-2,8-dihydroxy-N,N,10,10-tetramethyl-15-oxo-N-[3-[(1-oxododecyl)amino]propyl]-, trichloride, 5-oxide (9CI) (CA INDEX NAME)

TITLE: Cyclopropane methonium compounds  
 AUTHOR(S): Burger, Alfred; Bedford, G. R.  
 CORPORATE SOURCE: Univ. of Virginia, Charlottesville  
 SOURCE: Journal of Medicinal Chemistry (1963), 6(4), 402-5  
 CODEN: JMCMAR; ISSN: 0022-2623  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Unavailable  
 AB In a study of the effect of limiting the flexibility of the chains of methonium compds. on the pharmacol. actions of certain stereoisomers, analogs of hexamethonium and succinylcholine carrying a cis or trans oriented cyclopropane ring in the center of the chain were synthesized. The geometric isomers of bis(trimethylammoniummethyl) cyclopropane-1,2-dicarboxylate and of the homologous cyclopropane-1,2-diacetate ester diiodides caused predominantly neuromuscular block and resembled succinylcholine. The geometric isomers of 1,2-bis(β-trimethylammoniummethyl)cyclopropane diiodide exerted primarily ganglionic blockade of the hexamethonium type. The trans isomer was the more potent in each case.  
 IT 97299-16-6, Ammonium, [1,2,3-cyclopropanetriyltris(methylene)]tris [trimethyl-iodide]  
 (cyclopropyl derivs.)  
 RN 97299-16-6 CAPLUS  
 CN [1,2,3-Cyclopropanetriyltris(methylene)]tris[trimethylammonium iodide] (7CI) (CA INDEX NAME)



● 3 I-

=> d 117 31-43 ibib abs hitstr

L17 ANSWER 31 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1962:479123 CAPLUS  
 DOCUMENT NUMBER: 57:79123  
 ORIGINAL REFERENCE NO.: 57:15747d-e  
 TITLE: Neuromuscular-blocking agents. IX. Short-acting linear N,N,N-trisonium esters  
 AUTHOR(S): Carey Macleod, Fiona; Lewis, J. J.; Stenlake, J. B.; Williams, W. D.  
 CORPORATE SOURCE: Univ. Glasgow, UK  
 SOURCE: Journal of Pharmacy and Pharmacology (1961), 13(Suppl.), 103T-106T  
 CODEN: JPPMAB; ISSN: 0022-3573  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Unavailable  
 AB cf. CA 54, 11981c; 56, 7932c. The short series of linear trisonium esters are synthesized and are compared with tubocurarine, and suxamethonium for neuromuscular-blocking properties, potency, and toxicity. All compds. synthesized are tubocurarine-like except the methonium derivative (I), which is a depolarizing agent. I (3.0 mg./kg.) causes an inhibition of

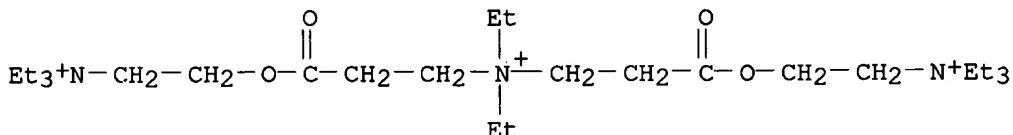
contractions and the development of contracture in the gastrocnemins musclesciatic nerve preparation of the pentobarbitone-anesthetized hen. Similar effects are obtained with 0.05 mg./kg. suxamethonium chloride.

IT 17089-56-4, Ammonium, bis(2-carboxyethyl)diethyl, iodide, diester with triethyl(2-hydroxyethyl)ammonium iodide 17089-57-5, Choline, iodide, diester with bis(2-carboxyethyl)ethylmethylammonium iodide

(nerve-muscle transmission blocking by)

RN 17089-56-4 CAPLUS

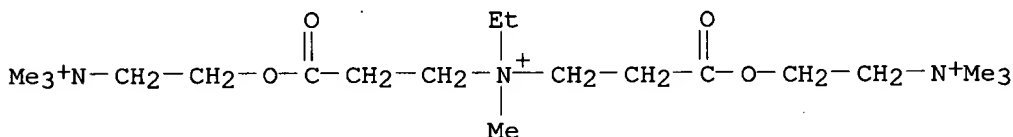
CN 1-Propanaminium, N,N-diethyl-3-oxo-N-[3-oxo-3-[2-(triethylammonio)ethoxy]propyl]-3-[2-(triethylammonio)ethoxy]-, triiodide (9CI) (CA INDEX NAME)



●3 I-

RN 17089-57-5 CAPLUS

CN 1-Propanaminium, N-ethyl-N-methyl-3-oxo-N-[3-oxo-3-[2-(trimethylammonio)ethoxy]propyl]-3-[2-(trimethylammonio)ethoxy]-, triiodide (9CI) (CA INDEX NAME)



●3 I-

L17 ANSWER 32 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1961:143790 CAPLUS

DOCUMENT NUMBER: 55:143790

ORIGINAL REFERENCE NO.: 55:27155e-i,27156a-i

TITLE: New class of local anesthetics.  
Hydroxyalkyliminobisacetamides

AUTHOR(S): Freed, Meier E.; Bruce, William F.; Hanslick, Roy S.;  
Maschitti, Albert

CORPORATE SOURCE: Wyeth Labs., Philadelphia, PA

SOURCE: Journal of Organic Chemistry (1961), 26,  
2378-83

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

AB cf. CA 53, 6088e. A series of hydroxyalkyliminobisacetamides, HOXN(CH<sub>2</sub>CONR<sub>1</sub>)CH<sub>2</sub>CONR<sub>2</sub>R<sub>3</sub> (where X is alkylene, cycloalkylene, or aralkylene, R, R<sub>1</sub>, R<sub>2</sub>, R<sub>3</sub> represent lower alkyl or aralkyl, and where R<sub>1</sub> may or may not equal R<sub>2</sub>R<sub>3</sub>), were prepared, examined for local anesthetic action, and studied for structure-activity relationships. The preparation of

all chloroacetamides, hydroxyalkylaminoacetamides, hydroxyalkyliminoacetamides and their esters were carried out essentially in the same manner.  $\text{PhCH}_2\text{CMe}_2\text{NHMe}$  (0.86 mole) in 500 ml.  $\text{PhMe}$  stirred 1 hr. at  $-15^\circ$  with addition of 0.40 mole  $\text{ClCH}_2\text{COCl}$ , the mixture filtered at  $20^\circ$ , the amine  $\text{HCl}$  salt washed with  $\text{PhMe}$ , the combined filtrate and washings dried, and the residue on evaporation distilled yielded 70.5%  $\text{PhCH}_2\text{CMe}_2\text{NMeCOCH}_2\text{Cl}$  (I),  $b_0.5\ 140-1^\circ$ .  $\text{HOCH}_2\text{CH}_2\text{NH}_2$  (0.1 mole) and 30 g. anhydrous powdered  $\text{Na}_2\text{CO}_3$  in 300 ml. well-stirred boiling  $\text{BuOH}$  slowly treated with 0.1 mole I in 50 ml.  $\text{BuOH}$ , the mixture refluxed 12 hrs., cooled, and filtered, and the residue on evaporation crystallized from  $\text{C}_6\text{H}_{14}$  yielded

63%  $\text{HOCH}_2\text{CH}_2\text{NHCH}_2\text{CONMeCMe}_2\text{CH}_2\text{Ph}$ , m.  $74.5-6.5^\circ$ ;  $\text{HCl}$  salt, m.  $163-4^\circ$ . Similarly were prepared and tabulated hydroxyalkylaminoacetamides,  $\text{RNHCH}_2\text{CONR}_1\text{Me}$  ( $\text{R, R}_1$ , and m.p.  $\text{HCl}$  salt given):  $\text{PhCHOHCH}_2$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $201-2^\circ$ ;  $\text{PhCHOHCMe}_2$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $189-90^\circ$ ;  $\text{HOCH}_2\text{CMe}_2$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $169-70^\circ$ ;  $(\text{HOCH}_2)_3\text{C}$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $175-6^\circ$ ;  $\text{HOCHMeCH}_2$ ,  $\text{PhCH}_2$ ,  $134-5^\circ$ . I (0.1 mole) and 20 g.  $\text{K}_2\text{CO}_3$  in 250 ml. boiling  $\text{BuOH}$  stirred with addition of 0.05 mole freshly distilled  $\text{HOCH}_2\text{CH}_2\text{NH}_2$ , the mixture refluxed 20 hrs. and the cooled mixture filtered, the filtrate washed (aqueous 5%  $\text{Na}_2\text{CO}_3$ ,  $\text{H}_2\text{O}$ ) and the dried ( $\text{MgSO}_4$ ) solution evaporated in  $\text{vacuo}$  yielded 71% hydroxyalkyliminobisacetamide,  $\text{RN}(\text{CH}_2\text{CONR}_1\text{R}_2)_2$  (II) ( $\text{R} = \text{HOCH}_2\text{CH}_2$ ,  $\text{R}_1 = \text{Me}$ ,  $\text{R}_2 = \text{PhCH}_2\text{CMe}_2$ ) (III), m.  $104-5^\circ$ ;  $\text{HCl}$  salt m.  $146-7^\circ$  ( $\text{MeOH}-\text{Me}_2\text{CO}$ ); nicotinic acid ester m.  $158-9^\circ$ . III (20 g.) in 100 ml. dry  $\text{CHCl}_3$  treated with 5 g.  $\text{SOC}_2$  in 25 ml.  $\text{CHCl}_3$ , the mixture stirred 3 hrs., and the residue on evaporation crystallized from alc.- $\text{Et}_2\text{O}$  yielded 79 g. II ( $\text{R} = \text{ClCH}_2\text{CH}_2$ ,  $\text{R}_1$

=  $\text{Me}$ ,

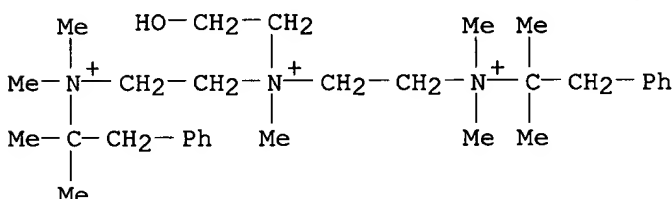
$\text{R}_2 = \text{PhCH}_2\text{CMe}_2$ )  $\text{HCl}$  salt (IV), m.  $155-6^\circ$  (alc.- $\text{Et}_2\text{O}$ ). IV (3 g.) in 20 ml.  $\text{MeOH}$  containing 3 g. anhydrous  $\text{NH}_3$  heated 18 hrs. at  $90^\circ$  in a pressure tube, the cooled mixture and  $\text{MeOH}$  rinsings filtered from  $\text{NH}_4\text{Cl}$ , freed from  $\text{MeOH}$  and excess  $\text{NH}_3$ , and taken up in 50 ml.  $\text{Me}_2\text{CHOH}$ , and the filtered solution treated with dry  $\text{HCl}$  and diluted with 150 ml. dry  $\text{Et}_2\text{O}$  yielded 40.5% II ( $\text{R} = \text{H}_2\text{NCH}_2\text{CH}_2$ ,  $\text{R}_1 = \text{Me}$ ,  $\text{R}_2 = \text{PhCH}_2\text{CMe}_2$ ), m.  $231-2^\circ$ . III (0.02 mole) in 150 ml. dry  $\text{Et}_2\text{O}$  added slowly with stirring to 1.8 g.  $\text{LiAlH}_4$  in 300 ml. dry  $\text{Et}_2\text{O}$ , the mixture refluxed 25 hrs. before cautious decomposition with 8 ml.  $\text{H}_2\text{O}$ , the dried  $\text{Et}_2\text{O}$  layer treated with  $\text{HCl}$ , the oily product triturated with  $\text{Me}_2\text{CO}$ , and the product (29.3%) recrystd. from  $\text{MeOHMe}_2\text{CO}$  yielded  $\text{HOCH}_2\text{CH}_2\text{N}(\text{CH}_2\text{CH}_2\text{NMe}_2\text{CH}_2\text{Ph})_2$ , m.  $229-30^\circ$  (decomposition); tri- $\text{HCl}$  salt m.  $239-40^\circ$ ;  $\text{MeI}$  salt, m.  $122-3^\circ$ ; tri- $\text{MeI}$  salt, m.  $154-5^\circ$ . To obtain the bis compds.

with sterically hindered amino alcs., the use of a higher boiling solvent (such as  $\text{PhOMe}$ ) was necessary. Phys. and pharmacol. data are tabulated for the various series of compds.,  $\text{RN}(\text{CH}_2\text{CONR}_1\text{R}_2)_2$  ( $\text{R, R}_1, \text{R}_2$ , b.p./mm., duration of activity on rabbit cornea and % solution given):  $\text{HOCH}_2\text{CH}_2$ , ( $\text{R}_1\text{R}_2=\text{CH}_2\text{CH}_2$ ,  $203-5^\circ/1.0$ , neg., 0.1;  $\text{HOCH}_2\text{CH}_2$ ,  $\text{Me}(\text{CH}_2)_3$ ,  $\text{Me}(\text{CH}_2)_3$ ,  $208-10^\circ/0.5$ , 25 min., 0.01;  $\text{MeCHOHCH}_2$ ,  $\text{Me}(\text{CH}_2)_3$ ,  $\text{Me}(\text{CH}_2)_3$ ,  $200-5^\circ/0.1$ , neg., 0.1;  $\text{HOCH}_2\text{CH}_2$ ,  $\text{Me}_2\text{CHCH}_2$ ,  $\text{Me}_2\text{CHCH}_2$ ,  $170-1^\circ/0.5$ , neg., 0.1;  $\text{HO}(\text{CH}_2)_3$ ,  $\text{Me}_2\text{CHCH}_2$ ,  $\text{Me}_2\text{CHCH}_2$ ,  $190-2^\circ/0.5$ , neg., 0.1;  $\text{HOCH}_2\text{CMe}_2$ ,  $\text{Me}(\text{CH}_2)_3$ ,  $\text{Me}(\text{CH}_2)_3$ ,  $155-60^\circ/0.5$ , neg., 0.1;  $\text{HOCH}_2\text{CMe}_2$ ,  $\text{MeCH}_2$ ,  $\text{Me}(\text{CH}_2)_3$ ,  $170-5^\circ/0.5$ , neg., 0.1;  $\text{HO}(\text{CH}_2)_{22}$ ,  $\text{Me}(\text{CH}_2)_4$ ,  $\text{Me}(\text{CH}_2)_4$ ,  $230-5^\circ/1.0$ , neg., 0.1;  $\text{HO}(\text{CH}_2)_2$ ,  $\text{C}_6\text{H}_{11}$ ,  $\text{C}_6\text{H}_{11}$ , - ( $\text{HCl}$  salt m.  $215-16^\circ$ ), neg., 0.1;  $\text{HO}(\text{CH}_2)_2$ ,  $\text{Me}(\text{CH}_2)_5$ ,  $\text{Me}(\text{CH}_2)_5$ ,  $194-6^\circ/0.5$ , 48 min., 0.1. For  $\text{RN}(\text{CH}_2\text{CONR}_1\text{R}_2)_2$  ( $\text{R, R}_1, \text{R}_2$ , m.p. of base or  $\text{HCl}$  salt (or b.p./mm.), duration in min. and % solution given):

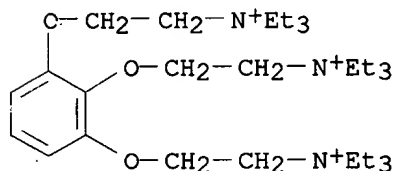
$\text{HO}(\text{CH}_2)_2$ ,  $\text{Me}$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $104-4.5^\circ$ , 25, 0.0005;  $\text{HOCHMeCH}_2$ ,  $\text{Me}$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $113-14^\circ$ , 28, 0.0001;  $\text{HOCH}_2\text{CHET}$ ,  $\text{Me}$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $144-5^\circ$  ( $\text{HCl}$  salt), 37, 0.0005;  $\text{HO}(\text{CH}_2)_3$ ,  $\text{Me}$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $164-5^\circ$  ( $\text{HCl}$  salt), 82, 0.1;  $\text{HO}(\text{CH}_2)_6$ ,  $\text{Me}$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $250-60^\circ/0.002$ , neg., 0.1;  $(\text{HOCH}_2)_3\text{C}$ ,  $\text{Me}$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $157-8^\circ$ , 24, 0.001; 2- $\text{HOC}_6\text{H}_{10}$ ,  $\text{Me}$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $108.0-8.5^\circ$ , 75, 0.0025;  $\text{PhCHOHCH}_2$ ,  $\text{Me}$ ,  $\text{PhCH}_2\text{CMe}_2$ ,  $182-3^\circ$  ( $\text{HCl}$  salt), 24, 0.001;  $\text{PhCHOHCMe}_2$ ,

Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 203-4° (HCl salt), neg., 0.1; HO(CH<sub>2</sub>)<sub>2</sub>, Me, C<sub>6</sub>H<sub>11</sub>, 190-5°/1.0, neg., 0.1; HO(CH<sub>2</sub>)<sub>2</sub>, H, PhCH<sub>2</sub>CH<sub>2</sub>, 72-3° (HCl salt), neg., 0.1; HO(CH<sub>2</sub>)<sub>2</sub>, Me(CH<sub>2</sub>)<sub>3</sub>, PhCH<sub>2</sub>, 118° (HCl salt), 29, 0.001; HOCHMeCH<sub>2</sub>, Me(CH<sub>2</sub>)<sub>5</sub>, PhCH<sub>2</sub>, 195-200°/0.05, 44, 0.1; HOCHMeCH<sub>2</sub>, H, 2,6-Me<sub>2</sub>C<sub>6</sub>H<sub>3</sub>, 193-4° (HCl salt), neg. 0.1. For RN(CH<sub>2</sub>CONR<sub>1</sub>R<sub>2</sub>)CH<sub>2</sub>CONR<sub>3</sub>R<sub>4</sub> (R, R<sub>1</sub>, R<sub>2</sub>, R<sub>3</sub>, R<sub>4</sub>, b.p./mm, or m.p. of base or HCl salt, duration, and % solution given): HO(CH<sub>2</sub>)<sub>2</sub>, MeCH<sub>2</sub>, MeCH<sub>2</sub>, Me(CH<sub>2</sub>)<sub>3</sub>, 203-5°/1.0, 29. 0.1; HO(CH<sub>2</sub>)<sub>2</sub>, Me(CH<sub>2</sub>)<sub>2</sub>, Me(CH<sub>2</sub>)<sub>2</sub>, Me(CH<sub>2</sub>)<sub>3</sub>, Me<sub>2</sub>CHCH<sub>2</sub>, 198-200°/0.5, 21, 0.1; HO(CH<sub>2</sub>)<sub>2</sub>, MeCH<sub>2</sub>, MeCH<sub>2</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 121-2°, neg., 0.1; HO(CH<sub>2</sub>)<sub>2</sub>, Me(CH<sub>2</sub>)<sub>4</sub>, Me(CH<sub>2</sub>)<sub>4</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 92-3°, 63, 0.1; HO(CH<sub>2</sub>)<sub>3</sub>, Me<sub>2</sub>CHCH<sub>2</sub>, Me<sub>2</sub>CHCH<sub>2</sub>, Me, C<sub>6</sub>H<sub>11</sub>, 205-80/1.0, neg., 0.1; HO(CH<sub>2</sub>)<sub>2</sub>, Me, PhCH<sub>2</sub>CHMe, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, hygroscopic, 55, 0.001; HO(CH<sub>2</sub>)<sub>2</sub>, H, PhCH<sub>2</sub>CH<sub>2</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 158° (HCl salt), neg., 0.1; HO(CH<sub>2</sub>)<sub>2</sub>, H, HOCH<sub>2</sub>CH<sub>2</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 42° (HCl salt), neg., 0.1; HO(CH<sub>2</sub>)<sub>2</sub>, H, Me(CH<sub>2</sub>)<sub>5</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 260°/1.0, 9, 0.05. For XCH<sub>2</sub>CH<sub>2</sub>N(CH<sub>2</sub>CONR<sub>1</sub>R<sub>2</sub>)<sub>2</sub> (X, R<sub>1</sub>, R<sub>2</sub>, m.p. HCl salt, duration, and % solution): MeCO<sub>2</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 169-70°, 32, 0.001; Me(CH<sub>2</sub>)<sub>10</sub>CO<sub>2</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 143-5°, 38, 0.01; p-MeC<sub>6</sub>H<sub>4</sub>CO<sub>2</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 168-9°, 42, 0.001; p-O<sub>2</sub>NC<sub>6</sub>H<sub>4</sub>CO<sub>2</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 168-9°, 27, 0.0005; MeCO<sub>2</sub>, Me(CH<sub>2</sub>)<sub>3</sub>, Me(CH<sub>2</sub>)<sub>3</sub>, 212-14°/0.05(base), 32, 0.01; m-C<sub>1</sub>C<sub>6</sub>H<sub>4</sub>CO<sub>2</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 87-8° (base, from Me<sub>2</sub>CHOH-petr. ether), active, 0.1; (3-C<sub>5</sub>H<sub>4</sub>N)CO<sub>2</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 158-9°, 35, 0.0005; p-MeOC<sub>6</sub>H<sub>4</sub>CO<sub>2</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 126-7°, active, 0.1; p-H<sub>2</sub>NC<sub>6</sub>H<sub>4</sub>CO<sub>2</sub>, Me, PhCH<sub>2</sub>CMe<sub>2</sub>, 199-200°, active, 0.1. Iminoacetamides in which the amido N was derived from aliphatic amines had relatively little local anesthetic action and were more toxic than those derived from aralkyl amines. The use of PhCH<sub>2</sub>CMe<sub>2</sub>NHMe produced the highest degree of local anesthetic activity in II. Substitution of PhCH<sub>2</sub>CHMeNHMe in 1 amide group halved the activity. In the alkanolamine moiety, use of a sterically hindered base (H<sub>2</sub>N<sup>+</sup>CMe<sub>2</sub>CH<sub>2</sub>OH) markedly reduced activity. Separation of HO from the tertiary amino group by interposition of CH<sub>2</sub> groups reduced activity. The activity of HOCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>N(CH<sub>2</sub>CONMeCMe<sub>2</sub>CH<sub>2</sub>Ph)<sub>2</sub> was 1/500 of that of the homologous HOCH<sub>2</sub>CH<sub>2</sub>N(CH<sub>2</sub>CONMeCMe<sub>2</sub>CH<sub>2</sub>Ph)<sub>2</sub>. Replacement of HO by NH<sub>2</sub> or Cl, and **quaternization** of the tertiary amine or reduction of the **amide** groups to tertiary amines all resulted in nearly complete loss of activity. The activity was not increased by **ester** formation.

IT 119722-18-8, Ammonium, [(2 hydroxyethylimino)diethylene]bis[( $\alpha$ , $\alpha$ -dimethylphenethyl)dimethyl- iodide], methiodide  
(preparation of)  
RN 119722-18-8 CAPLUS  
CN [(2-Hydroxyethylimino)diethylene]bis[( $\alpha$ , $\alpha$ -dimethylphenethyl)dimethylammonium iodide] methiodide (6CI) (CA INDEX NAME)



DOCUMENT NUMBER: 54:126132  
 ORIGINAL REFERENCE NO.: 54:24016f-g  
 TITLE: Action of some **quaternary** ammonium salts  
 with curare-like effect on the polarographic behavior  
 of cystine  
 AUTHOR(S): Serban, Mihail  
 SOURCE: Acad. rep. populare Romine, Inst. biochim., Studii  
 cercetari biochim. (1958), 1, 369-79  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Unavailable  
 AB The action of d-tuocurarine, flaxedyl, decamethonium, succinylcholine,  
 and "C100" on the polarog. wave of cystine was studied. All of these  
 substances increased the catalytic current, i.e. the height of the cystine  
 wave, the magnitude of the effect depending on the concentration of the salts.  
 IT 65-29-2, Ammonium, [v-phenenyltris(oxyethylene)]tris[triethyl-  
 iodide]  
 (cystine polarog. in presence of)  
 RN 65-29-2 CAPLUS  
 CN Ethanaminium, 2,2',2'''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-,  
 triiodide (9CI) (CA INDEX NAME)



● 3 I<sup>-</sup>

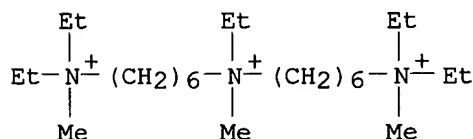
L17 ANSWER 34 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1960:62276 CAPLUS  
 DOCUMENT NUMBER: 54:62276  
 ORIGINAL REFERENCE NO.: 54:11981c-h  
 TITLE: Neuromuscular blocking agents. IV. Synthesis and study  
 of N- and S-alkyl variants of dihexasulfonium and  
 dihexazonium triethiodides  
 AUTHOR(S): Carey, Fiona M.; Edwards, D.; Lewis, J. J.; Stenlake,  
 J. B.  
 SOURCE: Journal of Pharmacy and Pharmacology (1959),  
 11, Suppl. 70T-86T  
 CODEN: JPPMAB; ISSN: 0022-3573  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Unavailable  
 AB cf. C.A. 53, 9032c. N,S,N- and N,N,N-Tri onium compds., related to  
 dihexasulfonium and dihexazonium in which N-alkyl substituents were  
 varied, were prepared N,S,N-Trionium compds. were prepared from either  
 bis(6-dimethylaminohexyl)sulfide or bis(6-diethylaminohexyl)sulfide by  
 refluxing with the alkyl halide in EtOH, evaporating, and crystallizing  
 Compds. with  
 min. of reflux, % yield, m.p., and crystallization solvent were:  
 7-ethyl-7-thioniatridecylenebis(dimethylammonium) triiodide, 35, 61,  
 137-7.5°, EtOH; 7-butyl-7-thioniatridecylenebis(dimethylbutylammonium) triiodide, 40, 51, 131-1.5°, EtOH-Me2COEt2O;  
 7-methyl-7-thioniatridecylenebis(diethylmethylammonium) triiodide, 20, 94,  
 135-6°, EtOH; 7-propyl-7-thioniatridecylenebis(diethylpropylammonium)

m) triiodide, 45, 52, 125.5-26°, EtOH-Et2O. Et  
 N,N-dipropyladipamate (from Et H adipate), yellow oil, b0.35  
 144-6°, nD22 1.4550, 87.6%. 6-Hydroxyhexyldipropylamine (from  
 N,N-dipropyladipate by LiAlH4) b0.65 115-17°, nD22 1.4533, 95%.  
 6-Propylaminohexyldipropylamine (from reflux of 6 -  
 hydroxyhexyldipropylamine and HBr, then propylamine), b0.45  
 115-17°, nD22, 1.4463, 70.6%. N,N-Dipropyladipamic acid (by  
 hydrolysis of Et N,N-dipropyladipamate in alc. KOH), yellow viscous oil,  
 b0.5 198°, nD25 1.4723, 91.9%. Bis(6-dipropylaminohexyl)propylamin  
 e (from reflux of N,N-dipropyladipamic acid in C6H6 and SOC12), pale  
 yellow oil, b0.65 211°, nD21 1.4582, 77.5%. N,N-Diethyladipamic  
 acid, yellow viscous oil, b0.5 182°, nD20.5 1.4733, 95.78.  
 Bis(6-diethylaminohexyl)ethylamine (from N,N-diethyladipamic acid and  
 excess 6-(diethylaminohexyl)ethylamine), pale yellow oil, b0.75  
 173-6°, nD25 1.4588, 55.9%. N,N,N-Trionium compds., prepared from  
 either bis(6-dipropylaminohexyl)propylamine or bis(6-  
 diethylaminohexyl)ethylamine by reflux with alkyl halide in EtOH, evaporation,  
 and crystallization, were (min. of reflux, % yield, m.p., and solvent given):  
 7-methyl-7-propyl-7-azoniatridecylenebis(dipropylmethylammonium)  
 triiodide, 10, 94, 239°, EtOH; 7-ethyl-7-propyl-7-  
 azoniatridecylenebis(dipropylethylammonium) triiodide, 35, 66,  
 221°, EtOH-Me2O-Et2O; 7,7-dipropyl-7-azoniatridecylenebis(tripropyl  
 ammonium) triiodide, 45, 12, 206-7°, Me2COEt2O;  
 7-ethyl-7-methyl-7-azoniatridecylenebis(diethylmethylammonium) triiodide,  
 5, 88, 227.5-8.5°, MeOH; 7-ethyl-7-propyl-7-  
 azoniatridecylenebis(diethylpropylammonium) triiodide, 30, 43,  
 220°, EtOH-Me2CO-Et2O; 7-ethyl-7-butyl-7-  
 azoniatridecylenebis(diethylpropylammonium) triiodide, 45, 61,  
 178°, Me2CO-Et2O. All the compds. tested qual. resembled  
 tubocurarine in their action. Stepwise replacement of Et by Me in  
 dihexasulfonium triethiodide (I) decreased potency. Potency also fell  
 when Et groups were replaced by Pr in I and dihexazonium triethiodide.

IT 1862-35-7, Ammonium, [(ethylimino)bis(hexamethylene)]bis[diethyl  
 methyl- iodide], methiodide 1862-36-8, Ammonium,  
 [(propylimino)bis(hexamethylene)]bis[diethylpropyl- iodide], ethiodide  
 1862-37-9, Ammonium, [(propylimino)bis(hexamethylene)]bis[ethyldip  
 royl- iodide], ethiodide 1862-38-0, Ammonium,  
 [(propylimino)bis(hexamethylene)]bis[tripropyl- iodide], propiodide  
 4055-56-5, Ammonium, [(propylimino)bis(hexamethylene)]bis[methyldi  
 propyl- iodide], methiodide 124245-58-5, Ammonium,  
 [(butylimino)bis(hexamethylene)]bis[diethylpropyl- iodide]-, ethiodide  
 (preparation of)

RN 1862-35-7 CAPLUS

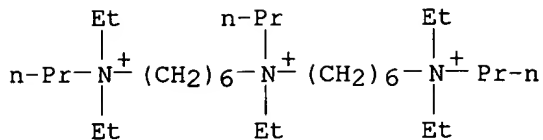
CN 1,6-Hexanediaminium, N-[6-(diethylmethylammonio)hexyl]-N,N',N'-triethyl-  
 N,N'-dimethyl-, triiodide (9CI) (CA INDEX NAME)



● 3 I-

RN 1862-36-8 CAPLUS

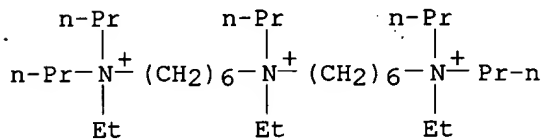
CN 1,6-Hexanediaminium, N-[6-(diethylpropylammonio)hexyl]-N,N',N'-triethyl-  
 N,N'-dipropyl-, triiodide (9CI) (CA INDEX NAME)



● 3 I<sup>-</sup>

RN 1862-37-9 CAPLUS

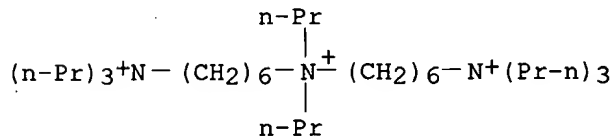
CN 1,6-Hexanediaminium, N,N'-diethyl-N-[6-(ethyldipropylammonio)hexyl]-N,N',N'-tripropyl-, triiodide (9CI) (CA INDEX NAME)



● 3 I<sup>-</sup>

RN 1862-38-0 CAPLUS

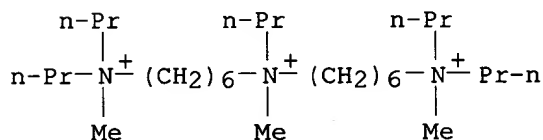
CN 1,6-Hexanediaminium, N,N,N,N',N'-pentapropyl-N'-(6-(tripropylammonio)hexyl)-, triiodide (9CI) (CA INDEX NAME)



● 3 I<sup>-</sup>

RN 4055-56-5 CAPLUS

CN Ammonium, [(methylpropyliminio)bis(hexamethylene)]bis[methyldipropyl-, triiodide (8CI) (CA INDEX NAME)]

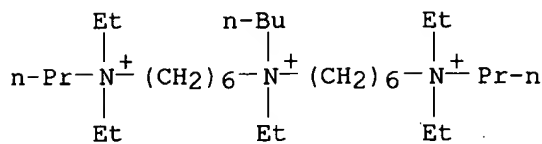


● 3 I<sup>-</sup>

RN 124245-58-5 CAPLUS

CN [(Butylimino)bis(hexamethylene)]bis[diethylpropylammonium iodide]

ethiodide (6CI) (CA INDEX NAME)



●3 I-

L17 ANSWER 35 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1960:39901 CAPLUS

DOCUMENT NUMBER: 54:39901

ORIGINAL REFERENCE NO.: 54:7895f-g

TITLE: Action of some **quaternary** ammonium base salts with curare-like activities upon the polarographic behavior of cystine

AUTHOR(S): Sherban, M.

SOURCE: Rev. chim. Acad. rep. populare Roumaine (1959), 4, 119-28

DOCUMENT TYPE: Journal

LANGUAGE: Russian

AB The following compds. with curare-like activity were investigated: d-tubocurarine, decamethonium, succinylcholine, flaxedil, and C100 (derivative of belladonna). The concentration of the curare-like compds. was kept at 10-5-10-4M while the concentration of cystine was maintained at 10-5M at pH

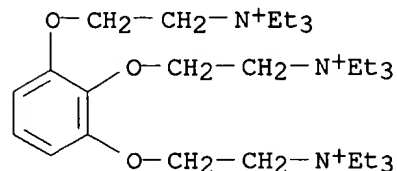
9.4.

The polarogram curves were traced at 3.8 v. and temperature 22°. Flaxedil was the most active. These **quaternary** compds. showed a pos. increase in the polarographic wave of cystine. The expts. indicated a close relation between the SH groups and the curare-like activities of compds. 13 references

IT 65-29-2, Ammonium, [v-phenenyltris(oxyethylene)]tris[triethyl-iodide]  
(effect on cystine polarography)

RN 65-29-2 CAPLUS

CN Ethanaminium, 2,2',2''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-, triiodide (9CI) (CA INDEX NAME)



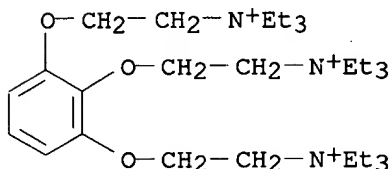
●3 I-

L17 ANSWER 36 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1959:119782 CAPLUS

DOCUMENT NUMBER: 53:119782

ORIGINAL REFERENCE NO.: 53:21361h-i,21362a  
 TITLE: Appearance of artifacts on chromatograms of  
**quaternary** ammonium compounds  
 AUTHOR(S): Crocker, Charity  
 CORPORATE SOURCE: Univ. Brazil, Rio de Janeiro  
 SOURCE: Journal of Chromatography (1959), 2, 115-16  
 CODEN: JOCRAM; ISSN: 0021-9673  
 DOCUMENT TYPE: Journal  
 LANGUAGE: English  
 AB Choline chloride, hexamethylenebis(carbamoylcholine iodide) (606H.C.), 1,4-bis(2-piperidinoethyl)piperazinedi-EtI (336H.C.), and gallamine triethiodide (Flaxedil), each containing residual  $CCl_3CO_2H$ , showed the presence of artifact spots when chromatographed on paper in alkaline solvents. The size of the artifact spot increased at the expense of the principal spot with increasing amts. of  $CCl_3CO_2H$ . Solvent mixts. used were EtOH-NH<sub>3</sub>, PrOH-NH<sub>3</sub>-H<sub>2</sub>O, and BuOH-C<sub>5</sub>H<sub>5</sub>N-H<sub>2</sub>O. When the artifact was eluted and rechromatographed in the same solvent or in acid solvents, it reappeared as such and not as the parent **quaternary** ammonium compound  
 IT 65-29-2, Ammonium, [v-phenenyltris(oxyethylene)]tris[triethyl-iodide]  
 (chromatography of, artifacts in)  
 RN 65-29-2 CAPLUS  
 CN Ethanaminium, 2,2',2''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-, triiodide (9CI) (CA INDEX NAME)



●3 I-

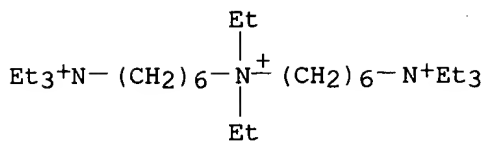
L17 ANSWER 37 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1959:50670 CAPLUS  
 DOCUMENT NUMBER: 53:50670  
 ORIGINAL REFERENCE NO.: 53:9032c-i,9033a  
 TITLE: Neuromuscular blocking agents. II. A series of  
 N,S,N-and N,N,N-trisethonium compounds  
 AUTHOR(S): Edwards, D.; Lewis, J. J.; Stenlake, J. B.; Zoha, M.  
 S.  
 SOURCE: Journal of Pharmacy and Pharmacology (1958),  
 10, 106T-121T  
 CODEN: JPPMAB; ISSN: 0022-3573  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Unavailable  
 AB cf. C.A. 52, 8036f. 6-Hydroxyhexyldiethylamine (35.2 g.) in 95 ml. 48% HBr and 33 ml. H<sub>2</sub>SO<sub>4</sub> was refluxed 4 hrs., cooled, poured into 1 l. H<sub>2</sub>O, Na<sub>2</sub>CO<sub>3</sub> added and the mixture extracted with CHCl<sub>3</sub>, and the dried extract (Na<sub>2</sub>SO<sub>4</sub>) evaporated in vacuo to obtain crude 6-bromohexyldiethylamine (I) as a reddish brown oil containing crystalline material. The I obtained and 40 ml. ethylamine refluxed 2 hrs. and EtNH<sub>2</sub> and CHCl<sub>3</sub> evaporated yielded a damp crystalline mass;

this basified, extracted with, Et2O, and the extract evaporated yielded 24.8 g. oil,  
distilled to yield 53% 6-ethylaminohexyldiethylamine (II), b0.55 86-9°, n17D 1.4493; di-HCl salt m. 172-3° (EtOH-Et2O).  
Crystalline material separated from crude II was  
1,1-diethyl-1-azacycloheptylinium bromide, m. 250° (decomposition) (EtOH-Et2O). II and 6-chlorohexyldiethylamine refluxed in xylene 5 hrs. formed 19% bis(6-diethylaminohexyl)ethylamine, pale yellow oil, b0.7 165-8°, n18D 1.4610; this was refluxed 10 min. with EtI to form 7,7-diethyl-7-azoniatridecylenebis(triethylammonium) triiodide, m. 261-2°, needles (EtOH). Bis(10-diethylaminodecyl) sulfide was refluxed with EtI to form 27% 11-ethyl-11-thoniaheneicosylenebis(triethylammonium) triiodide, m. 123.5-24°, needles (Me2CO-Et2O).  
10-Bromodecyldiethylamine (84%), b0.5 130°, n14D 1.4717, was obtained as an oil by the method for I. 10-Ethylaminodecyldiethylamine (76%), an oil, b0.8 133-5°, n18D 1.4535; di-HCl salt m. 147-8° (EtOH-Et2O). Bis(10-diethylaminodecyl)ethylamine (26.5%), pale yellow oil, b0.25 212-16°, n14D 1.4660; tri-HCl salt, m. 118° (Me2CO-Et2O). 11,11-Diethyl-11-azoniaheneicosylenebis(triethylammonium) triiodide (90%) m. 202.5-3.5° (Me2CO-Et2O).  
1,1-Bis(ethoxycarbonyl)-7-diethylaminoheptane (47.65%), pale yellow oil, b0.8 147-55°, n15.5D 1.4472, was used to prepare 62% Et 8-diethylaminocaprylate (III), an oil, b0.65 111-14°, n18D 1.4428; this reacted with EtI to form 7-ethoxycarbonylheptyltriethylammonium iodide, m. 64.5-5.5° (Me2CO-Et2O). III was reduced with LiAlH4 to obtain 90% 8-hydroxyoctyldiethylamine, an oil, b0.7 114-17°, n16.5D 1.4590; HCl salt m. 90-1° (EtOH-Et2O). 8-Chlorooctyldiethylamine (96%), an oil, b0.55 94-6°, n17D 1.4550 (literature, b11 130.5°, n18D 1.4535). Bis(8-diethylaminoctyl) sulfide (72%), straw-colored liquid, b0.65 210-12°, n18.5D 1.4768; di-HCl salt m. 145° (EtOH). 9-Ethyl-9-thoniaheptadecylenebis(triethylammonium) triiodide (47%) m. 159-60° (decomposition) (EtOH-Et2O).  
8-Ethylaminoctyldiethylamine (76%), an oil, b0.7 104-6°, n17.5D 1.4530; di-HCl salt, hygroscopic, m. 159.5-60.5° (EtOH-Et2O). Bis(8-diethylaminoctyl)ethylamine (18%), yellow oil, b0.8 230-50°, n17D 1.4642; tri-HCl salt m. 165-6° (decomposition) (Me2CO-Et2O).  
9,9-Diethyl-9-azoniaheptadecylenebis(triethylammonium) triiodide m. 251-2° (decomposition) (EtOH). 7-Dioxothiatridecylenebis(triethylammonium iodide) (47%), pale buff solid, m. 144-5° (Me2CO-Et2O). All the compds. tested showed neuromuscular blocking activity. Dihexazonium triethiodide and the sulfone 7-dioxothiatridecylenebis(triethylammonium iodide) (dihexone) showed tubocurarine-like activity; dioctasulfonium and dioctazonium triethiodides were predominantly tubocurarine-like but had some transitional properties. Didecasulfonium and didecazonium triethiodides resembled decamethonium. Dihexazonium triethiodide was equipotent with tubocurarine on the cat. Marked species variations in potency were noted.

IT 3756-18-1, Ammonium, [(ethylimino)bis(hexamethylene)]bis[triethyl-iodide], ethiodide 15159-46-3, Ammonium, [(ethylimino)bis(octamethylene)]bis[triethyl- iodide], ethiodide 102031-41-4, Ammonium, decamethylenebis[(10-diethylaminodecyl)diethyl- iodide], diethiodide 106715-64-4, Ammonium, hexamethylenebis[(6-diethylaminohexyl)diethyl- iodide], diethiodide 108019-73-4, Ammonium, [(ethylimino)bis(decamethylene)]bis[triethyl- iodide], ethiodide  
(preparation of)

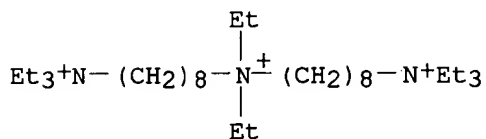
RN 3756-18-1 CAPLUS

CN 1,6-Hexanediaminium, N,N,N,N',N'-pentaethyl-N'-(6-(triethylammonio)hexyl)-, triiodide (9CI) (CA INDEX NAME)



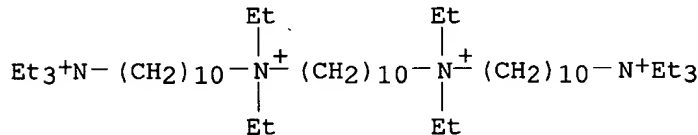
● 3 I<sup>-</sup>

RN 15159-46-3 CAPLUS  
 CN 1,8-Octanediaminium, N,N,N,N',N'-pentaethyl-N'-[8-(triethylammonio)octyl]-, triiodide (9CI) (CA INDEX NAME)



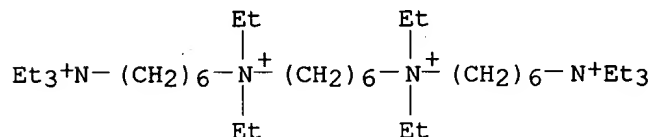
● 3 I<sup>-</sup>

RN 102031-41-4 CAPLUS  
 CN 3,3,14,14,25,25,36,36-Octaethyl-3,14,25,36-tetraazoniaoctatriacontane tetraiodide (6CI, 7CI) (CA INDEX NAME)



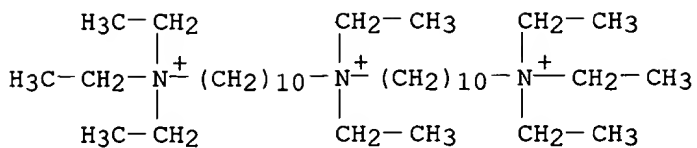
● 4 I<sup>-</sup>

RN 106715-64-4 CAPLUS  
 CN 3,3,10,10,17,17,24,24-Octaethyl-3,10,17,24-tetraazoniahexacosane tetraiodide (6CI, 7CI) (CA INDEX NAME)



● 4 I<sup>-</sup>

RN 108019-73-4 CAPLUS  
 CN 3,3,14,14,25,25-Hexaethyl-3,14,25-triaazoniaheptacosane triiodide (7CI) (CA INDEX NAME)



● 3 I-

L17 ANSWER 38 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1958:89882 CAPLUS

DOCUMENT NUMBER: 52:89882

ORIGINAL REFERENCE NO.: 52:15830a-g

TITLE: Isolation, characterization, and determination of basic organic active substances of various medicinals with disulfimides. I

AUTHOR(S): Runge, F.; Engelbrecht, H. J.; Franke, H.

CORPORATE SOURCE: Univ. Halle, Saale, Germany

SOURCE: Pharmazie (1957), 12, 8-13

CODEN: PHARAT; ISSN: 0031-7144

DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

AB cf. C.A. 50, 7803h. The disulfimides (I) as strong acids form with organic bases crystalline salts poorly soluble in water. The most useful I is (4-ClC<sub>6</sub>H<sub>4</sub>SO<sub>2</sub>)<sub>2</sub>NNa (II), prepared by mixing NH<sub>4</sub>Cl and p-ClC<sub>6</sub>H<sub>4</sub>SO<sub>2</sub>Cl in Me<sub>2</sub>CO, adding 30% NaOH solution, and heating gently to distil off the Me<sub>2</sub>CO. The base (IIA) is prepared by dissolving II in hot water and precipitating with HCl, dissolving in anhydrous Et<sub>2</sub>O, evaporating, and crystallizing from C<sub>6</sub>H<sub>6</sub>. IIA, m.

205-6°, is well suited for the isolation, characterization, and determination of primary, secondary, tertiary, and **quaternary** amines, preferable often to picric acid or perchlorates, and especially for the **quaternary** compds., with which they form stable crystalline compds. with sharp m.ps. The amine or its salt combines with I or their alkaline salts in an ionic reaction. The free base in Et<sub>2</sub>O solution may be left to react with I directly, or I Na is mixed in aqueous, alc., or Me<sub>2</sub>CO solution with

the **quaternary** base or its salts, advantageously with heat. An excess of either component retards crystallization, hence molar proportions must

be used as closely as possible. The following medicinals were crystallized (compound and m.p. of product with IIA (uncorrected) given): anesthesine 150-1°; procaine 141-2°; 2-diethylaminoethanol 76-7°; coramine 153-4°; dilatol [1-(p-hydroxyphenyl)-2-(1-methyl-3-phenylpropylamino)propanol-HCl] 175-6°; sympathol 168-9°; dispasmol (N-benzyl-N',N'-dimethyl-N-phenylethylenediamine) 141-2°; rodismim (N-benzyl-N',N'-diethyl-N-phenylethylenediamine) 94-5°, 2-( $\alpha$ -phenyl-o-tolyloxy)triethylamine-HCl 128-9°; aminopyrine 167-8°; megaphen from 50° (unsharp); sulfanilamide 197-8°; sulfapyridine 165-6°; sulfacetamide Na, yellow crystals 167-8°; sulfaguanidine 108-9°; elkosin 179-80°; nicotinic acid hydrazide from 206° (decompose); tetramethyldecamethylenediamine-di-MeBr 150-1°; flaxedil 62-4°; choline chloride succinate 78-80°; hexamethonium bromide 214-15°; benzedrine 182-3°; thiamine-HCl 206-7°; niacinamide 212-14°; 3-pyridyl benzyl carbonate,

weak rose-colored needles, 167-8°; atrophan, yellowish needles, 171-2°; urotropine 162-3°; ephedrine 184-5°; atropine 145-6°; hyoscyamine 123-5°; papaverine 118-19° (unsharp); hydrastinine 139-40°; and yohimbine 196-7°. To determine satisfactory methods for gravimetric analysis, various I salts of 2 medicinals were prepared. Thus, for casantin, were prepared compds. with (4-H<sub>2</sub>NC<sub>6</sub>H<sub>4</sub>SO<sub>2</sub>)<sub>2</sub>NH (m. 202-3°), II (m. 111-12°), and (3,4-C<sub>12</sub>C<sub>6</sub>H<sub>3</sub>SO<sub>2</sub>)<sub>2</sub>NH (III) (m. 146-7°). The last compound was found best because it had the lowest solubility, was well crystallized, and had a high

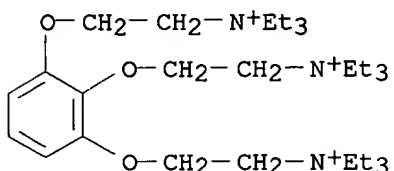
mol. weight. For hexamethonium bromide, salts were prepared with II (m. 214-15°) and with III (m. 207-8°). Gravimetric detns. for both compds. were more accurate than volumetric (argentimetric) detns. 11 references.

IT 65-29-2, Ammonium, [v-phenenyltris(oxyethylene)]tris[triethyl-iodide]

(detection of, and preparation of its salt with 4,4'-dichlorodibenzenesulfonamide)

RN 65-29-2 CAPLUS

CN Ethanaminium, 2,2',2'''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-, triiodide (9CI) (CA INDEX NAME)



●3 I-

L17 ANSWER 39 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1958:78488 CAPLUS

DOCUMENT NUMBER: 52:78488

ORIGINAL REFERENCE NO.: 52:13971a-c

TITLE: Influence on the metabolism of the eggs of *Psammechinus microtuberculatus* of **quaternary** ammonium compounds and phenothiazine derivatives

Hofmann, H.

AUTHOR(S): Friedrich Schiller Univ., Jena, Germany

CORPORATE SOURCE: Pharmazeutische Zentralhalle fuer Deutschland (1957), 96, 421-31

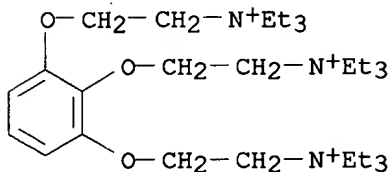
SOURCE: CODEN: PHZEAD; ISSN: 0369-9773

DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

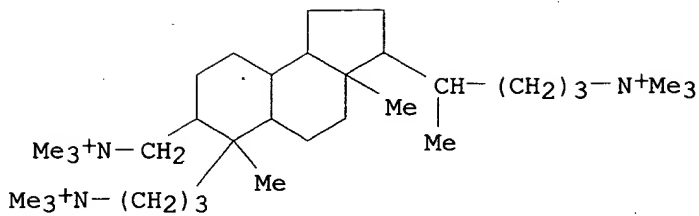
AB **Quaternary** ammonium compds. (d-tubocurarine chloride, Flaxedil, hexamethonium, decamethonium, and pendiomid) decreased the O consumption of fertilized and unfertilized eggs of *R. microtuberculatus*. The decrease in metabolism was inversely proportional to drug concentration. Phenothiazine derivs. of the chlorpromazine type decreased the O consumption of sea-urchin eggs; above a certain limiting concentration the decrease in metabolism rose sharply and led to complete inhibition of oxidation. When combined with ethylurethan (I) the **quaternary** ammonium compds. showed an additive effect. The combination of phenothiazine derivs. and I gave a potentiating action. By this means, a further difference of the ganglioplegic drugs from the phenothiazine derivs. has been found.

IT 65-29-2, Ammonium, [v-phenenyltris(oxyethylene)]tris[triethyl-  
iodide]  
(inhibition of sea-urchin egg metabolism by)  
RN 65-29-2 CAPLUS  
CN Ethanaminium, 2,2',2''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-,  
triiodide (9CI) (CA INDEX NAME)



●3 I<sup>-</sup>

L17 ANSWER 40 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
ACCESSION NUMBER: 1958:50718 CAPLUS  
DOCUMENT NUMBER: 52:50718  
ORIGINAL REFERENCE NO.: 52:9173i, 9174a-c  
TITLE: Multivalent **quaternary** ammonium compounds.  
VI. Some reaction products of bile acids and sterols  
AUTHOR(S): Lettre, H.; Gottstein, W.; Scholtissek, Ch.  
CORPORATE SOURCE: Univ. Heidelberg, Germany  
SOURCE: Monatshefte fuer Chemie (1957), 88, 715-20  
CODEN: MOCMB7; ISSN: 0026-9247  
DOCUMENT TYPE: Journal  
LANGUAGE: Unavailable  
AB cf. C.A. 51, 4409a. Some N derivs. of lithiobilanic acid (I) and  
sitosterol are prepared. I is treated with Ac<sub>2</sub>O followed by Me<sub>2</sub>NH to yield  
lithiobilanic acid 3-monodimethylamide, m. 251-2°. I with Ac<sub>2</sub>O  
followed by PC15 and then with Me<sub>2</sub>NH in Et<sub>2</sub>O gives an Et<sub>2</sub>O phase containing  
60%-70% I 3,4,24-tris(dimethylamide), m. 151-2°, purified by  
chromatography on Al<sub>2</sub>O<sub>3</sub>. The aqueous phase of the reaction yields 15-20% of I  
3,24-bis(dimethylamide) (II), m. 232-3°. II is esterified with  
CH<sub>2</sub>N<sub>2</sub> and reduced with LiAlH<sub>4</sub> in tetrahydrofuran to 3,4-secocholan-4-ol-  
3,24-bis(dimethylamine hydrochloride), m. 292-5° (decomposition). II is  
similarly reduced to 90% 3,4-secocholane-3,4,24-tris(dimethylamine  
hydrochloride), decompose 275°, which forms 3,4-secocholane-3,24-  
tris(trimethylammonium iodide), m. 290° (decomposition). The  
dicarboxylic acid of sitosterol (III), heated 2 hrs. with Ac<sub>2</sub>O gives 76%  
2,3-secositostanol-2,3-dicarboxylic acid anhydride, m. 176°. III  
di-Me **ester** is reduced by LiAlH<sub>4</sub> to 88% 2,3-secositostane-2,3-  
diol, m. 182-3° (MeOH). III with PC15 and Me<sub>2</sub>NH yields by  
chromatography on Al<sub>2</sub>O<sub>3</sub> 48% 2,3-secositostane-2,3-dicarboxylic acid  
dimethylamide, m. 106-7°, reduced by LiAlH<sub>4</sub> to 68%  
2,3-secositostane-2,3-bis(dimethylamine hydrochloride), m. 326°  
(decomposition). This compound with MeI gives 2,3-secositostane-2,3-  
bis(trimethylammonium iodide), m. 323°.  
IT 122387-46-6, 3,4-Secocholane-3,4,24-triamine, N<sub>3</sub>,N<sub>3</sub>,N<sub>4</sub>,N<sub>4</sub>,N<sub>24</sub>,N<sub>24</sub>-  
hexamethyl-, trimethiodide  
(preparation of)  
RN 122387-46-6 CAPLUS  
CN 3,4-Secocholane-3,4,24-triamine, N<sub>3</sub>,N<sub>3</sub>,N<sub>4</sub>,N<sub>4</sub>,N<sub>24</sub>,N<sub>24</sub>-hexamethyl-,  
trimethiodide (6CI) (CA INDEX NAME)



●3 I-

L17 ANSWER 41 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1957:6726 CAPLUS

DOCUMENT NUMBER: 51:6726

ORIGINAL REFERENCE NO.: 51:1456g-i

TITLE: Antagonists to the neuromuscular block produced by Sarin

AUTHOR(S): Kunkel, A. M.; Wills, J. H.; Monier, J. S.

CORPORATE SOURCE: Army Chem. Center, MD

SOURCE: Proceedings of the Society for Experimental Biology and Medicine (1956), 92, 529-32

CODEN: PSEBAA; ISSN: 0037-9727

DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

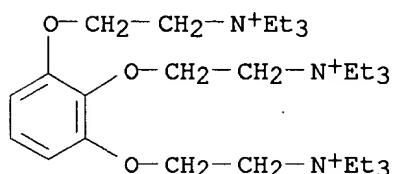
AB Large i.v. doses of Sarin decrease the twitch height of the cat gastrocnemius-soleus muscle group excited by maximal elec. stimulation of the sciatic nerve at 2-s. intervals. Various compds. containing **quaternary** N atoms, including several atropine derivs., overcome the decrease in twitch height. Some compds. with significant anticholinesterase activity enhance the Sarin-induced decrease in twitch height despite the abolition by Sarin of demonstrable cholinesterase activity in the muscle.

IT 65-29-2, Ammonium, [v-phenenyltris(oxyethylene)]tris[triethyl-iodide]

(as antagonist for Sarin)

RN 65-29-2 CAPLUS

CN Ethanaminium, 2,2',2'''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-, triiodide (9CI) (CA INDEX NAME)



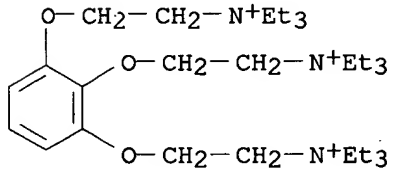
●3 I-

L17 ANSWER 42 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1952:6411 CAPLUS

DOCUMENT NUMBER: 46:6411

ORIGINAL REFERENCE NO.: 46:1150b-d  
 TITLE: Synthetic curarizing agents. III. Succinylcholine and its aliphatic derivatives  
 AUTHOR(S): Bovet, D.; Bovet-Nitti, F.; Guarino, S.; Longo, V. G.; Fusco, R.  
 CORPORATE SOURCE: Ist. super. sanità, Rome  
 SOURCE: Archives Internationales de Pharmacodynamie et de Therapie (1951), 88, 1-50  
 CODEN: AIPTAK; ISSN: 0003-9780  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Unavailable  
 AB cf. C.A. 44, 1603c. Succinylcholine diiodide produces typical curarization in frogs and mammals. Birds, and the isolated frog rectus abdominis, show a nicotine-like contracture. In the dog, there is no effect on the blood pressure or cardiac rhythm except in large doses when the drug causes hypertension and tachycardia. Excess salivation and bronchial secretion occur. The direct excitability of the gastrocnemius muscle is not affected. In the series  $X(R)3N(CH_2)nOOC(CH_2)mCOO(CH_2)nN(R)3$   $X$  the curarizing action is most marked if a chain of about 10 C and O atoms separate the **quaternary** nitrogens, and the substituent groups on the N are Me. Some of the series  $I(CH_3)3N(CH_2)5COOCH_2CH_2N(CH_3)3$   $I$ , and the series  $I(CH_3)3NCH_2CH_2OOC(CH_2)2COOCH_2CH_2N(CH_3)3$   $I$  are also active, but branching of the chain or the introduction of a third or fourth **quaternary** N destroys the activity.  
 IT 65-29-2, Ammonium, [v-phenenyltris(oxyethylene)]tris[triethyl-iodide]  
     (pharmacology of)  
 RN 65-29-2 CAPLUS  
 CN Ethanaminium, 2,2',2'''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-, triiodide (9CI) (CA INDEX NAME)



●3 I-

L17 ANSWER 43 OF 43 CAPLUS COPYRIGHT 2005 ACS on STN  
 ACCESSION NUMBER: 1948:1255 CAPLUS  
 DOCUMENT NUMBER: 42:1255  
 ORIGINAL REFERENCE NO.: 42:274a-g  
 TITLE: Curarizing properties of phenolic ethers with **quaternary** ammonium groups  
 AUTHOR(S): Bovet, Daniel; Depierre, France; de Lestrange, Yvonne  
 SOURCE: Compt. rend. (1947), 225, 74-6  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Unavailable  
 AB A study was made of the action on striated muscle of synthetic curarizing agents consisting of ethers formed from choline and homologous amino alcs. with phenols and polyphenols. The following compds. were used:  
 $C_6H_5OCH_2CH_2N(CH_3)3I$  (I),  $1,3-C_6H_4[OCH_2CH_2N(CH_3)3I]_2$  (II),  
 $C_6H_5OCH_2N(C_2H_5)3I$  (III),  $1,2-C_6H_4[OCH_2CH_2N(C_2H_5)3I]_2$  (IV),  
 $1,3-C_6H_4[OCH_2CH_2N(C_2H_5)3I]_2$  (V),  $1,4-C_6H_4[OCH_2CH_2N(C_2H_5)I_3I]_2$  (VI),

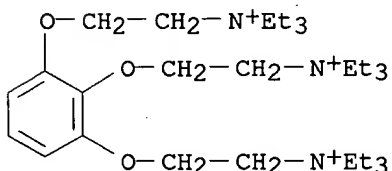
1,2,3-C<sub>6</sub>H<sub>3</sub>[OCH<sub>2</sub>CH<sub>2</sub>N(C<sub>2</sub>H<sub>5</sub>)<sub>3</sub>I]3 (VII). In the choline series, I and II were effective in the rabbit in doses of 4 mg. per kg. given intravenously. In the triethylcholine series, III has very slight activity, but IV, V, VI, and especially VII, are very active, the last causing curarization of 3 hrs. duration in the rabbit, in a dose of 0.7 mg. per kg. In the **quaternary** amines, choline and the **ester** salts of choline, particularly acetylcholine and butyl- $\beta$ -ethylcholine (Dale, C.A. 9, 104) the curarizing effects seem to be closely connected with other cholinergic, muscarinic, and nicotinic manifestations of the mol. It was observed that in the ethers of the polyphenols studied, the cardiovascular effects are considerably attenuated. While I causes hypertension comparable to that produced by nicotine, II exerts only a weak nicotinic action. Likewise, the hypotensive and cardiomoderating effect of III is considerably weakened by the introduction of 1 or 2 more **quaternary** ammonium groups. The effects of VII are particularly striking. This compound is very active in the frog, which is immobilized by it in doses of 10 mg. per kg. In the mouse, the toxic doses are 5.5, 15, and 425 mg., given intravenously, subcutaneously, and per os. In the rabbit, the toxic dose is 0.7 mg. given intravenously, and 2-3.5 mg. per kg. subcutaneously. This represents about 5 times the activity of tubocurarine. In a rabbit given artificial respiration, total paralysis lasts 2.5 hrs. with a dose of 7 mg. and 6 hrs. after 35 mg. To kill a rabbit under these conditions, 350 mg., or about 500 times the toxic dose is necessary. In the chloralosed dog given 0 by tracheal catheter, the response of the gastrocnemius muscle to elec. excitation at the peripheral end of the sciatic nerve decreases in amplitude, then disappears at the same time as paralysis of the respiratory muscles occurs. The vagus is paralyzed, presumably at the synapses of its cardiac ganglia, but acetylcholine still exerts a large part of its normal effect. The injection of eserine (1-2 mg. per kg. in the atropinized dog) or of prostigmine, results in rapid recovery of muscular excitability. Injection of a dose of VII sufficient to cause curarization for several hrs. has no effect on blood pressure. In this respect the compound is superior to natural curare.

IT 65-29-2, Ammonium, [v-phenenyltris(oxyethylene)]tris[triethyl-iodide]

(curarizing action of)

RN 65-29-2 CAPLUS

CN Ethanaminium, 2,2',2''-[1,2,3-benzenetriyltris(oxy)]tris[N,N,N-triethyl-, triiodide (9CI) (CA INDEX NAME)



● 3 I -

⇒